PATENT APPLICATION TRANSMITTAL LETTER

To the Commissioner of Patents and Trademarks:

CERTIFICATE OF MAILING

I hereby certify that this correspondence is being deposited with the United States Postal Service as "Express Mail" in an envelope addressed to: Assistant Commissioner for Patents, Box Patent Application, Washington, D.C. 20231, Express Mail Receipt No. EL.253288301US

Nuly 1999

Kay Speake



Transmitted	herewith for filing	under 35 U.S.C. 1	11 and 37 C.F	.R §1.53 is	the patent	application o	f:
Ronald Joh	nston Hill and Gan	y Noel Hannan					
entitled:"	Genetic Sequences	Encoding Steroic	l and Juvenile	Hormone	Receptor	Polypeptides	and

Enclosed are:

- X 131 pages of written description, claims, abstract and sequence listing.
- X 4 sheets of drawings.

Insecticidal Modalities Therefor"

- an assignment of the invention to
- <u>X</u> unexecuted declaration of the inventor and power of attorney.
- __ a certified copy of a _____ application.
- _ associate power of attorney.
- a verified statement to establish small entity status under 37 CFR §1.9 and §1.27.
- information disclosure statement.
- __ preliminary amendment
- other:

CLAIMS AS FILED

	Number Filed	Number Extra	Rate	Fee
BASIC FEE	,	, , ,	\$760	\$760
TOTAL CLAIMS	39-20=	19	x \$18	\$342
INDEPENDENT CLAIMS	13-3=	10	x \$78	\$780
MULTIPLE DEPENDENT CLAIM PRESENT			x \$260	

* Number extra must be zero or larger

TOTAL \$1882

If applicant has small entity status under 37 C.F.R.

1.9 and 1.27, then divide total fee by 2, and enter amount here.

SMALL ENTITY TOTAL

\$

<u>X</u> - -	No filing fee is enclosed at this time. A check in the amount of \$ to cover the filing fee is enclosed. The Commissioner is hereby authorized to charge and credit Deposit Account No. 07-1969 as described below. A duplicate copy of this sheet is enclosed.
	Charge the amount of \$ as filing fee. Credit any overpayment. Charge any additional filing fees required under 37 CFR 1.16 and 1.17. Charge the issue fee set in 37 CFR 1.18 at the mailing of the Notice of Allowance, pursuant to 37 CFR 1.311(b). Other
X	Benefit of Prior U.S. Application(s) (35 USC 120)
	Applicant claims priority under 35 USC 120 to the following application(s):
	PCT application PCT/AU99/00033 filed 15 January 1999
	Benefit of Prior U.S. Provisional Application(s) (35 USC 119(e))
	Applicant claims priority under 35 USC 120 to the following application(s):
<u>X</u>	Benefit of Prior Foreign Application(s) (35 USC 119)
	Applicant claims priority under 35 USC 119 to the following application:
	Australian application PP1356/98 filed 15 January 1998

Donna M. Ferber Reg. No. 33,878

GREENLEE WINNER and SULLIVAN, P.C.

5370 Manhattan Circle, Suite 201

Boulder, CO 80303 Phone: (303) 499-8080 Fax: (303) 499-8089

email: winner@greenwin.com ks: July 1, 1999

ks: July 1, 1999 Docket 53-99

APPLICATION FOR LETTERS PATENT

Inventors:

Ronald Johnston Hill Garry Noel Hannan

GENETIC SEQUENCES ENCODING STEROID AND JUVENILE HORMONE RECEPTOR POLYPEPTIDES AND INSECTICIDAL MODALITIES THEREFOR

CERTIFICATE OF MAILING

I hereby certify that this correspondence is being deposited with the United States Postal Service as "Express Mail" in an envelope addressed to: Assistant Commissioner for Patents, Box Patent Application, Washington, D.C. 20231, Express Mail Receipt No. EL253288301US

Prepared by:

GREENLEE, WINNER and SULLIVAN, P.C. 5370 Manhattan Circle Suite 201 Boulder, Colorado 80303 (303) 499-8080

30

35

5

10

GENETIC SEQUENCES ENCODING STEROID AND JUVENILE HORMONE RECEPTOR POLYPEPTIDES AND INSECTICIDAL MODALITIES THEREFOR

This application is a continuation-in-part of Australian application PP1356/98 filed 15 January 1998 and PCT application PCT/AU99/00033 filed 15 January 1999, both incorporated herein by reference to the extent that there is no inconsistency with the present disclosure.

FIELD OF THE INVENTION

The present invention relates generally to novel genetic sequences encoding receptor polypeptides and insecticidal modalities therefor, which insecticidal modalities are based upon non-polypeptide insect hormones and their receptors. More specifically, the present invention provides isolated nucleic acid molecules encoding polypeptides comprising functional steroid hormone and juvenile hormone receptors, in particular isolated nucleic acid molecules which encode polypeptides comprising the Lucilia cuprina and Myzus persicae ecdysone receptors and juvenile hormone receptors. In a particularly preferred embodiment, the present invention relates to isolated nucleic acid molecules which encode the L. cuprina and M. persicae EcR polypeptide subunits and EcR partner protein (USP polypeptide) subunits which form functional heterodimeric ecdysone receptor, and to the L. cuprina and M. persicae USP polypeptide of the juvenile hormone receptor. The present invention further relates to the production of functional recombinant insect receptors and recombinant polypeptide subunits thereof and derivatives and analogues thereof. The present invention further relates to the uses of the recombinant receptor and isolated nucleic acid molecules of the present invention in the regulation of gene expression. The present invention further relates to screening systems and methods of identifying insecticidally-active agents which are capable of agonising or antagonising insect receptor function, such as molecules and/or ligands which associate with steroid receptors or juvenile hormone receptors so as to modify the affinity of said receptors for their cognate cis-acting response elements (e.g., insect steroid response elements, juvenile hormone response elements) in the genes which they regulate, or alternatively or in addition, which modify the affinity of said receptors for their cellular stimuli (e.g., insect steroids or juvenile hormones) or analogues thereof, or alternatively or in addition, which act as insecticides by virtue of their ability to agonise or antagonise the activity of insect hormones, such as by mimicry of a ligand which binds to said receptor or a ligand-binding region thereof. The invention further extends to such compounds and/or ligands.

This specification contains nucleotide and amino acid sequence information prepared using the program Patentin Version 2.0, presented herein after the bibliography. Each nucleotide

-2-

or amino acid sequence is identified in the sequence listing by the numeric indicator <210> followed by the sequence identifier (e.g. <210>1, <210>2, etc). The length, type of sequence (DNA, protein (PRT), etc) and source organism for each nucleotide or amino acid sequence are indicated by information provided in the numeric indicator fields <211>, <212> and <213>, 5 respectively. Nucleotide and amino acid sequences referred to in the specification are defined by the information provided in numeric indicator field <400> followed by the sequence identifier (eg. <400>1, <400>2, etc).

The designation of nucleotide residues referred to herein are those recommended by the 10 IUPAC-IUB Biochemical Nomenclature Commission, wherein A represents Adenine, C represents Cytosine, G represents Guanine, T represents thymine, Y represents a pyrimidine residue, R represents a purine residue, M represents Adenine or Cytosine, K represents Guanine or Thymine, S represents Guanine or Cytosine, W represents Adenine or Thymine, H represents a nucleotide other than Guanine, B represents a nucleotide other than Adenine, 15 V represents a nucleotide other than Thymine, D represents a nucleotide other than Cytosine and N represents any nucleotide residue.

Bibliographic details of the publications referred to in this specification are collected at the end of the description.

20

As used herein the term "derived from" shall be taken to indicate that a specified integer may be obtained from a particular source albeit not necessarily directly from that source.

Throughout this specification, unless the context requires otherwise, the word "comprise", or 25 variations such as "comprises" or "comprising", will be understood to imply the inclusion of a stated step or element or integer or group of steps or elements or integers but not the exclusion of any other step or element or integer or group of elements or integers.

Those skilled in the art will appreciate that the invention described herein is susceptible to 30 variations and modifications other than those specifically described. It is to be understood that the invention includes all such variations and modifications. The invention also includes all of the steps, features, compositions and compounds referred to or indicated in this specification,

- 3 -

individually or collectively, and any and all combinations or any two or more of said steps or features.

The present invention is not to be limited in scope by the specific embodiments described 5 herein, which are intended for the purposes of exemplification only. Functionally-equivalent products, compositions and methods are clearly within the scope of the invention, as described herein.

BACKGROUND TO THE INVENTION

10 International Patent Application No WO91/13167 (applicant, The Board of Trustees of Leyland Stanford University, and hereinafter referred to as WO91/13167) describes the identification, characterization, expression and uses of insect steroid receptors and DNA sequences encoding same and, in particular, the identification, characterization, expression and uses of the steroid receptor of the common fruit fly, Drosophila melanogaster.

15

It has been found by the present inventors that the limited homology between the D. melanogaster steroid receptor-encoding gene sequences and the steroid receptor-encoding sequences derived from other insects, in particular those derived from diptera such as the Australian sheep blowfly L. cuprina, hemiptera such as aphids, scale insects and leaf hoppers, 20 coleoptera, neuroptera, lepidoptera, and ants, as well as from helminths and protozoa, prevents the routine isolation of DNA sequences encoding steroid receptors and/or juvenile hormone receptors from these latter-mentioned organisms.

Moreover, the present inventors have discovered that the D. melanogaster steroid receptor 25 described in WO91/13167 is temperature-sensitive, showing reduced activity at temperatures above 30°C, such as at temperatures about 37°C, particularly at low concentrations of the receptor. Accordingly, the D. melanogaster steroid receptor described in WO91/13167 is of limited utility at physiological temperatures applicable to animal or bacterial cells. Moreover, wherein it is desirable to produce a biologically-active steroid receptor using in vivo or in situ 30 expression systems, which expression systems routinely utilise cells or tissues in the temperature range of about 28°C to about 42°C, the D. melanogaster steroid receptor is also

-4-

of limited utility.

In work leading up to the present invention, the present inventors developed a novel screening protocol, involving the utilisation of highly-degenerate oligonucleotide probes and primers 5 derived from the amino acid sequences of the DNA-binding domains of the D. melanogaster and Chironomus tentans ecdysone receptor polypeptides, to identify nucleotide sequences encoding novel steroid receptor polypeptides and novel insect juvenile hormone receptor polypeptides. The present inventors have further identified specific regions within these novel polypeptides which are suitable for use in preparing a surprising range of novel steroid 10° receptor polypeptide derivatives and insect juvenile hormone receptor polypeptide derivatives. The novel steroid receptor polypeptides and novel insect juvenile hormone receptor polypeptides of the present invention, and derivative polypeptides thereof, and assembled steroid receptors and insect juvenile hormone receptors derived from said polypeptides and derivatives, and nucleic acid molecules encoding same as exemplified herein, provide the 15 means for developing a wide range of insecticidally-active agents, as well as methods for the regulated production of bioactive molecules. In particular, the present invention provides the means for developing specific ligands which bind to and either agonise or antagonise the steroid receptors and/or juvenile hormone receptors and/or polypeptide subunits thereof as described herein, thereby functioning as highly-specific insecticides, offering significant 20 commercial and environmental benefits.

The present inventors have been surprisingly successful in characterizing the ecdysone receptor and juvenile hormone receptor derived from insects of the orders Diptera and Hemiptera, and polypeptide components thereof and functional derivatives of said polypeptides and receptors, particularly in light of the extreme difficulties in dealing with these organisms. The nature of these molecules was unknown prior to the present invention.

The various aspects of this invention overcome the problems associated with *Drosophila* ecdysone receptors which lack thermal stability. Moreover, those aspects of the invention pertaining to methods of screening for insecticidally active agents do not involve competition assays which are generally complex, and often inaccurate or difficult to calibrate.

- 5 -

SUMMARY OF THE INVENTION

One aspect of the present invention provides an isolated nucleic acid molecule comprising a nucleotide sequence which encodes or is complementary to a sequence which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide or a bioactive derivative or analogue thereof, wherein said polypeptide:

- (i) is selected from the list comprising EcR polypeptide of an steroid receptor, the partner protein (USP polypeptide) of an steroid receptor and the USP polypeptide of a juvenile hormone receptor; and
- (ii) comprises an amino acid sequence that is at least 40% identical to any one of the amino acid sequences set forth in <400>2, <400>4, <400>6, <400>10, <400>12 or <400>14.

In an alternative embodiment, the present invention provides an isolated nucleic acid molecule comprising a nucleotide sequence which encodes or is complementary to a sequence which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide or a bioactive derivative or analogue thereof, wherein said polypeptide:

- (i) is selected from the list comprising EcR polypeptide of an steroid receptor, the partner protein (USP polypeptide) of an steroid receptor and the USP polypeptide of a juvenile hormone receptor; and
- (ii) comprises an amino acid sequence that is at least 40% identical to any one of the amino acid sequences encoded by the plasmids deposited under any one of AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568.

In a further alternative embodiment, the isolated nucleic acid molecule of the present invention comprises a nucleotide sequence which encodes or is complementary to a sequence which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide or a bioactive derivative or analogue thereof, wherein said nucleotide sequence is selected from the list comprising:

(i) a nucleotide sequence having at least 40% identity to any one of the nucleotide sequences set forth in <400>1, <400>3, <400>5, <400>9, <400>11 or <400>13 or a complementary nucleotide sequence thereto;

15

P:\OPER\MRO\ECDYSONE.CIP - 1/7/99

- (ii) a nucleotide sequence that is capable of hybridising under at least low stringency conditions to any one of the nucleotide sequences set forth in <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11 or <400>13 or to a complementary nucleotide sequence thereto;
- 5 (iii) a nucleotide sequence that is capable of hybridising under at least low stringency conditions to a nucleotide sequence contained in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568.;
 - (iv) a nucleotide sequence that is at least 40% identical to a nucleotide sequence contained in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568; and
 - (v) a nucleotide sequence that is amplifiable by PCR using a nucleic acid primer sequence set forth in any one of <400>15, <400>16, <400>17, <400>18, <400>19 or <400>20.

In a further alternative embodiment, the present invention provides an isolated nucleic acid molecule which encodes a steroid receptor polypeptide and comprises the nucleotide sequence set forth in <400>1 or a complementary nucleotide sequence thereto.

- In a further alternative embodiment, the present invention provides an isolated nucleic acid molecule which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide and comprises the nucleotide sequence set forth in <400>3 or <400>13 or a complementary nucleotide sequence thereto.
- 25 In a further alternative embodiment, the present invention provides an isolated nucleic acid molecule which encodes a steroid receptor polypeptide and comprises the nucleotide sequence set forth in <400>5 or <400>7 or <400>8 or <400>9 or a complementary nucleotide sequence thereto.
- 30 In a further alternative embodiment, the present invention provides an isolated nucleic acid molecule which encodes a steroid receptor polypeptide or a juvenile hormone receptor

15

20

30

P.\OPER\MRO\ECDYSONE.CIF - 1/7/99

polypeptide and comprises the nucleotide sequence set forth in <400>11 or a complementary nucleotide sequence thereto.

A second aspect of the present invention provides a method of identifying an isolated nucleic acid molecule which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide comprising the steps of:

- (i) hybridising genomic DNA, mRNA or cDNA with a hybridisation-effective amount of one or more probes selected from the list comprising:
 - (a) probes comprising at least 10 contiguous nucleotides in length derived from any one of <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11, <400>13, <400>15, <400>16, <400>17, <400>18, <400>19 or <400>20 or a complementary nucleotide sequence thereto:
 - (b) probes comprising at least 10 contiguous nucleotides in length derived from a cDNA contained in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568; and
 - (c) hybridisation probes comprising the nucleotide sequences set forth in any one of <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11, or <400>13 or a complementary nucleotide sequence thereto or a homologue, analogue or derivative thereof which is at least 40% identical to said sequence or complement; and
- (ii) detecting the hybridisation,

In an alternative embodiment, the inventive method comprises the steps of:

- (i) annealing to genomic DNA, mRNA or cDNA, one or more PCR primers comprising at least 10 contiguous nucleotides in length derived from the group consisting of:
 - (a) a primer derived from any one of <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11, <400>13, <400>15, <400>16, <400>17, <400>18, <400>19 or <400>20 or a complementary nucleotide sequence thereto; and
 - (b) a primer derived from a cDNA contained in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566,

10

15

20

25

30

P:\OPER\MRO\ECDYSONE.CIP - 171/99

- 8 -

NM99/04567, or NM99/04568; and

- (ii) amplifying a nucleotide sequence which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide in a polymerase chain reaction.
- 5 In a further alternative embodiment, the inventive method comprises the steps of:
 - (i) amplifying a nucleotide sequence which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide in a polymerase chain reaction using one or more PCR primers comprising at least 10 contiguous nucleotides in length from the group consisting of:
 - (a) a primer derived from any one of <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11, <400>13, <400>15, <400>16, <400>17, <400>18, <400>19 or <400>20 or a complementary nucleotide sequence thereto; and
 - (b) a primer derived from a cDNA contained in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568;
 - (ii) hybridising the amplified nucleotide sequence to genomic DNA, mRNA or cDNA with a hybridisation-effective amount of one or more probes selected from the group consisting of:
 - (a) a probe comprising at least 10 contiguous nucleotides in length derived from any one of <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11, <400>13, <400>15, <400>16, <400>17, <400>18, <400>19 or <400>20 or a complementary nucleotide sequence thereto:
 - (b) a probe comprising at least 10 contiguous nucleotides in length derived from a cDNA contained in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568; and
 - (c) hybridisation probes comprising the nucleotide sequences set forth in any one of <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11, or <400>13 or a complementary nucleotide sequence thereto or a homologue, analogue or derivative thereof which is at least 40% identical to said sequence or complement; and
 - (iii) detecting the hybridisation.

P'\OPER\MRO\ECDYSONE.CIP - 1/7/99

A third aspect of the present invention provides a genetic construct comprising the subject isolated nucleic acid molecule which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide, operably linked to a promoter sequence. Preferably, the subject nucleic acid molecule is in an expressible format, such that it is possible to produce a recombinant polypeptide therefrom.

Accordingly, a fourth aspect of the invention provides a recombinant or isolated polypeptide comprising a steroid receptor polypeptide or juvenile hormone receptor polypeptide or a bioactive derivative or analogue thereof, wherein said polypeptide:

- (i) is selected from the list comprising EcR polypeptide of a steroid receptor, the partner protein (USP polypeptide) of a steroid receptor and the USP polypeptide of a juvenile hormone receptor; and
- (ii) comprises an amino acid sequence that is at least 40% identical to any one of the amino acid sequences set forth in <400>2, <400>4, <400>6, <400>10, <400>12 or <400>14;

wherein said polypeptide is substantially free of naturally-associated insect cell components.

In an alternative embodiment, the invention provides a recombinant or isolated polypeptide comprising a steroid receptor polypeptide or juvenile hormone receptor polypeptide or a bioactive derivative or analogue thereof, wherein said polypeptide:

- (i) is selected from the list comprising EcR polypeptide of a steroid receptor, the partner protein (USP polypeptide) of a steroid receptor and the USP polypeptide of a juvenile hormone receptor; and
- (ii) comprises an amino acid sequence that is at least 40% identical to a polypeptide encoded by the cDNA present in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568;

wherein said polypeptide is substantially free of naturally-associated insect cell components.

30 A fifth aspect of the invention provides a cell comprising the subject isolated nucleic acid molecule which encodes a steroid receptor polypeptide or a juvenile hormone receptor

- 10 -

polypeptide.

5

25

In a preferred embodiment, the cell of the present invention expresses the polypeptide encoded by the nucleic acid molecule.

In a preferred embodiment, the cell expresses a steroid receptor polypeptide or a fragment thereof which receptor is capable of binding to an insect steroid or analogue thereof or a candidate insecticidally active agent to form an activated complex, and comprises a nucleic acid sequence encoding a bioactive molecule or a reporter molecule operably linked to one or more insect steroid response elements which on binding of the said activated complex promotes transcription of the nucleic acid sequence, wherein said cell on exposure to insect steroid or an analogue thereof, regulates expression of said bioactive molecule or allows detection of said reporter molecule.

15 In a further aspect of this invention, there is provided an animal (such as a mammal), microorganism, plant or aquatic organism, containing one or more cells as mentioned above.

A further aspect of the present invention provides a method of identifying a modulator of insect steroid receptor-mediated gene expression or insect juvenile hormone receptor-mediated gene expression comprising:

- (i) assaying the expression of a reporter gene in the presence of a recombinant or isolated insect steroid receptor polypeptide or a juvenile hormone receptor polypeptide of the invention and a potential modulator; and
- (ii) assaying the expression of a reporter gene in the presence of a recombinant or isolated insect steroid receptor polypeptide or a juvenile hormone receptor polypeptide of the invention and without said potential modulator; and
 - (ii) comparing expression of the reporter gene in the presence of the potential modulator to the expression of a reporter gene in the absence of the potential modulator.
- 30 wherein said reporter gene is placed operably under the control of a steroid response element (SRE) to which said insect steroid receptor binds or a promoter sequence comprising said

- 11 -

SRE.

10

15

A still further aspect of the invention provides a method of identifying a potential insecticidal compound comprising:

- 5 (i) assaying the binding directly or indirectly of a recombinant or isolated insect steroid receptor polypeptide or a juvenile hormone receptor polypeptide of the invention to a steroid response element (SRE) to which said insect steroid receptor binds, in the presence of a candidate compound; and
 - (ii) assaying the binding directly or indirectly of a recombinant or isolated insect steroid receptor polypeptide or a juvenile hormone receptor polypeptide of the invention to a steroid response element (SRE) to which said insect steroid receptor binds, in the absence of said candidate compound; and
 - (ii) comparing the binding assayed at (i) and (ii), wherein a difference in the level of binding indicates that the candidate compound possesses potential insecticidal activity.

A still further aspect of the invention provides a method of identifying a candidate insecticidally-active agent comprising the steps of:

- a) expressing an EcR polypeptide of a steroid receptor or a fragment thereof 20 which includes the ligand-binding region, optionally in association with an EcR partner protein (USP polypeptide) of a steroid receptor or ligand binding domain thereof, optionally in association with an insect steroid or analogue thereof so as to form a complex;
 - b) purifying or precipitating the complex;
- c) determining the three-dimensional structure of the ligand binding domain of the complex; and
 - d) identifying compounds which bind to or associate with the three-dimensional structure of the ligand binding domain, wherein said compounds represent candidate insecticidally-active agents.

30

A still further aspect of the invention provides a method of identifying a candidate insecticidally-

P:\OPER\MRO\ECDYSONE,CIP - 1/7/99

active agent comprising the steps of:

- expressing a USP polypeptide of a juvenile hormone receptor or a fragment thereof which includes the ligand-binding region, optionally in association with an EcR polypeptide of a steroid receptor or ligand binding domain thereof, and optionally in association with an insect steroid or analogue thereof, so as to form a complex;
- b) purifying or precipitating the complex;
- c) determining the three-dimensional structure of the ligand binding domain of the complex; and
- identifying compounds which bind to or associate with the three-dimensional d) 10 structure of the ligand binding domain, wherein said compounds represent candidate insecticidally-active agents.

In another aspect this invention relates to a method or assay for screening insecticidally active compounds which comprises reacting a candidate insecticidal compound with a steroid 15 receptor polypeptide or fragment thereof encompassing the ligand binding domain, or complex thereof with a partner protein or a fragment thereof which encompasses the ligand binding domain, and detecting binding or absence of binding of said compound so as to determine insecticidal activity.

- 20 A still further aspect of the invention provides a synthetic compound which interacts with the three dimensional structure of a polypeptide or protein selected from the list comprising:
 - an EcR polypeptide of a steroid receptor or a fragment thereof; (i)
 - (ii) an EcR partner protein (USP polypeptide) of a steroid receptor or a fragment thereof;
- 25 (iii) a USP polypeptide of a juvenile hormone receptor; and
 - (iv) a functional receptor or protein complex formed by association of (i) and (ii), wherein said compound is capable of binding to said polypeptide or protein to agonise or antagonise the binding activity or bioactivity thereof.
- 30 Preferably, the synthetic compounds are derived from the three dimensional structure of insect steroid receptor(s) or juvenile hormone receptor(s) which compounds bind to said receptor(s)

- 13 -

and have the effect of either inactivating the receptor(s) or potentiating the activity of the receptor(s). More preferably, the compounds mimic the three-dimensional structure of a ligand which binds to the receptor(s) and more preferably, mimic the three-dimensional structure of a ligand which binds to the ligand-binding region of said receptor(s).

5

In a still further aspect of this invention, there is provided a screening system for insecticidally active agents comprising a nucleotide sequence encoding a steroid receptor or a fragment thereof, and a nucleotide sequence encoding a partner protein or a fragment thereof which associates with the receptor so as to confer enhanced affinity for insect steroid response elements, enhanced affinity for insect steroids or analogues thereof, or insecticidally active agents and/or thermostability or enhanced thermostability of said receptor, which receptor and partner protein is capable of binding to a candidate insecticidally active agent to form an activated complex, and a nucleic acid sequence encoding a bioactive molecule or a reporter molecule operably linked to one or more insect steroid response elements which on binding of the said activated complex regulates transcription of the nucleic acid sequence, wherein on exposure to said agent expression of the bioactive molecule or reporter molecule correlates with insecticidal activity.

In another aspect of this invention, there is provided a method for the regulated production of a bioactive molecule or a reporter molecule in a cell, said method comprising the steps of introducing into said cell:

- a) a nucleotide sequence encoding a steroid receptor or a fragment thereof which is capable of binding an insect steroid or analogue thereof, to form an activated complex; and
- b) a nucleotide sequence encoding said bioactive molecule or reporter molecule operably linked to one or more insect steroid response elements which on binding of the said activated complex regulates transcription of the nucleic acid sequence encoding said bioactive molecule or reporter molecule,

wherein exposing the cell to an insect steroid or analogue thereof regulates expression of the bioactive molecule or reporter molecule.

<400>3:

10

25

P:\OPER\MRO\ECDYSONE.CIP - 1/1/99

- 14 -

SUMMARY OF SEQUENCE LISTING

<400>1: The nucleotide sequence of the open reading frame of a cDNA molecule which encodes the EcR polypeptide subunit of the L. cuprina ecdysone receptor and amino acid sequence therefor.

5 **<400>2:** The amino acid sequence of the EcR polypeptide subunit of the *L. cuprina* ecdysone receptor.

The nucleotide sequence of the open reading frame of a cDNA molecule which encodes the EcR partner protein (USP polypeptide) subunit of the *L. cuprina* ecdysone receptor and/or which encodes the USP polypeptide subunit of the *L. cuprina* juvenile hormone receptor, and amino acid sequence therefor.

<400>4: The amino acid sequence of the EcR partner protein (USP polypeptide) subunit of the L. cuprina ecdysone receptor and/or the amino acid sequence of the USP polypeptide subunit of the L. cuprina juvenile hormone receptor.

7400>5: The nucleotide sequence of a cDNA molecule which encodes part of the EcR polypeptide subunit of the M. persicae ecdysone receptor and amino acid sequence therefor.

<400>6: The amino acid sequence of a part of the EcR polypeptide subunit of the M. persicae ecdysone receptor.

7: The nucleotide sequence of the EcR probe 1 which is specific for genetic sequences encoding the EcR polypeptide subunit of aphid ecdysone receptors, in particular the EcR polypeptide subunit of the M. persicae ecdysone receptor.

<400>8: The nucleotide sequence of the EcR probe 2 sequence which is specific for genetic sequences encoding the EcR polypeptide subunit of aphid ecdysone receptors, in particular the EcR polypeptide subunit of the *M. persicae* ecdysone receptor.

<400>9: The nucleotide sequence of the open reading frame of a cDNA molecule which encodes the EcR polypeptide subunit of the *M. persicae* ecdysone receptor and amino acid sequence therefor.

<400>10: The amino acid sequence of the EcR polypeptide subunit of the M. persicae ecdysone receptor.

<400>11: The nucleotide sequence of the open reading frame of a cDNA molecule which

15

<400>16:

PAOPERIMRO/ECDYSONE.CIP - 1/1/99

- 15 -

encodes the EcR partner protein (USP polypeptide) subunit of the M. persicae ecdysone receptor and/or which encodes the USP polypeptide subunit of the M. persicae juvenile hormone receptor, and amino acid sequence therefor.

<400>12: The amino acid sequence of the EcR partner protein (USP polypeptide) subunit 5 of the M. persicae ecdysone receptor and/or the amino acid sequence of the USP polypeptide subunit of the M. persicae juvenile hormone receptor.

<400>13: The nucleotide sequence of a 150 base-pair probe which is specific for genetic sequences encoding the EcR partner protein (USP polypeptide) subunit of L. cuprina ecdysone receptor and/or the USP polypeptide subunit of the L. cuprina juvenile hormone receptor, and amino acid sequence therefor.

<400>14: The amino acid sequence encoded by the nucleotide sequence of <400>13, comprising amino acid residues 108-149 of the EcR partner protein (USP polypeptide) subunit of the L. cuprina ecdysone receptor and/or amino acid residues 108-149 of the amino acid sequence of the USP polypeptide subunit of the L_ cuprina juvenile hormone receptor set forth herein as <400>4.

<400>15: The nucleotide sequence of the degenerate primer Rdna3.

The nucleotide sequence of the degenerate primer Rdna4. <400>17: The nucleotide sequence of the primer Mdna1.

The nucleotide sequence of the primer Mdna2. <400>18:

20 <400>19: The nucleotide sequence of the primer AP1.

<400>20: The nucleotide sequence of the degenerate primer AP2.

BRIEF DESCRIPTION OF THE DRAWINGS

- 25 Figure 1 is a graphical representation showing function of the EcR polypeptide subunit of the L. cuprina ecdysone receptor in vivo. CHO cells were cotransfected with:
 - one of the following expression plasmids: pSGDmEcR, pSGLcEcR, or the (1) parental expression plasmid pSG5 as a control, at 1µg/ml;
 - (2) plasmid p(EcRE),-CAT (1 µg/ml); and
- 30 (3) an independent reporter plasmid, pPGKLacZ, at 1 µg/ml. CAT expression was induced with Muristerone A at either 10 µM or 50 µM while control cells

received only the carrier ethanol. ELISA kits were used to quantify the synthesis of CAT and β-galactosidase in extracts of cells forty eight hours after transfection. The level of CAT was normalized to the level of β-galactosidase in the same extract. Fold-induction represents the normalized values for CAT gene expression in cells transfected with pSGDmEcR, pSGLcEcR or pSG5 in the presence of hormone, relative to the normalized values for CAT gene expression in cells transfected with the same plasmid, but in the absence of hormone. The average values of three independent experiments are shown and the error bars indicate standard error of the mean.

- Figure 2 is a copy of a graphical representation showing the activity of plasmid pSGLD and pSGDL, containing chimeric EcR polypeptide subunits of insect ecdysone receptors, produced as described in the Examples. Cotransfection assays were performed as described in the Examples using plasmids pSGLD and pSGDL and the CAT reporter plasmid p(EcRE)₇ -CAT (1ug/ml) and an independent reporter, pPGKLacZ at 1 µg/ml each. CAT/b-Gal (%) refers to CAT reporter activity expressed as a percentage relative to β-galactosidase activity produced by the internal control reporter, pPGKLacZ.
- Figure 3 is a copy of a graphical representation showing the binding activity in extracts of Sf9 and Sf21 cells containing a baculvirus expressing LcEcRDEF and LcUSPDEF, as described in the Examples. Control cells contained baculovirus expressing β-galactosidase and CAT only.
- Figure 4 is a graphical representation showing the ecdysteroid binding activities of an *in vitro*-translated *Myzus persicae* EcR (MpEcR) polypeptide, an *in vitro*-translated *Myzus persicae* 25 USP (MpUSP) polypeptide, and an *in vitro*-translated complex of the *M.persicae* EcR and USP polypeptides.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

One aspect of the present invention provides an isolated nucleic acid molecule comprising a nucleotide sequence which encodes or is complementary to a sequence which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide or a bioactive

- 17 -

derivative or analogue thereof, wherein said polypeptide:

- (i) is selected from the list comprising EcR polypeptide of a steroid receptor, the partner protein (USP polypeptide) of a steroid receptor and the USP polypeptide of a juvenile hormone receptor; and
- 5 (ii) comprises an amino acid sequence that is at least 40% identical to any one of the amino acid sequences set forth in <400>2, <400>4, <400>6, <400>10, <400>12 or <400>14.

Accordingly, the isolated nucleic acid molecule of the invention may comprise a fragment of a nucleotide sequence encoding a full-length receptor polypeptide.

It is to be understood that a "fragment" of a nucleotide sequence encoding an EcR polypeptide subunit of a steroid receptor or an EcR partner protein (USP polypeptide) of a steroid receptor or a USP polypeptide of a juvenile hormone receptor, refers to a nucleotide sequence encoding a part or fragment of such a receptor which is capable of binding or associating with an insect steroid or an analogue thereof, or a candidate insecticidally active compound. Fragments of a nucleotide sequence would generally comprise in excess of twenty contiguous nucleotides derived from the base sequence and may encode one or more domains of a functional insect steroid receptor or juvenile hormone receptor.

20

Preferably, the isolated nucleic acid molecule of the invention encodes an ecdysteroid receptor polypeptide. Those skilled in the art are aware that ecdysteroid receptors derived from insects are heterodimeric receptors comprising an EcR polypeptide subunit and an EcR partner protein (USP polypeptide) (see also Jones and Sharp, 1997). In this regard, the present inventors have discovered that the USP polypeptide of the insect juvenile hormone receptor is structurally-identical to the EcR partner protein of the ecdysteroid receptor of the present invention, however juvenile hormone receptors comprise monomers or multimers of the USP polypeptide acting without the EcR polypeptide subunit that is present in the ecdysteroid receptors. Accordingly, the present invention extends equally to nucleotide sequences encoding polypeptides of both the ecdysteroid receptors and polypeptides of the juvenile hormone receptors of insects.

ű

10

25

P;\OPER\MRO\ECDYSONE.CIP - 1/7/99

- 18 -

More preferably, the isolated nucleic acid molecule of the invention encodes an ecdysteroid receptor that is modulated by one or more of the steroids ecdysone, ponasterone A, or muristerone, or an analogue of an ecdysteroid.

5 The isolated nucleic acid molecule of the invention may be derived from any organism that contains steroid receptors that are responsive to ecdysteroids or ecdysteroid-like compounds or juvenile hormones, or analogues of such receptor-ligands. Accordingly, the present invention is not to be limited in any of its embodiments to the particular source of the subject nucleic acid, or polypeptide encoded therefor.

Preferably, the isolated nucleic acid molecule of the invention is derived from insects,

helminths (nematodes, cestodes, trematodes), protozoa, and ants, amongst others.

More preferably, the isolated nucleic acid molecule of the invention is derived from an insect selected from the list comprising diptera, hemiptera, coleoptera, neuroptera, lepitdoptera and ants, amongst others. Still more preferably, the isolated nucleic acid molecule of the present invention is derived from aphids, scale insects, leaf hoppers, white fly, and blowflies such as sheep blowflies.

The present invention does not extend to amino acid sequences comprising the complete EcR polypeptide subunit of the *D. melanogaster* ecdysone receptor as described in WO91/13167. However, this exclusion is made on the understanding that the present invention does encompass chimeric genes and fusion proteins which include the *D. melanogaster* nucleotide and amino acid sequences, respectively.

In a particularly preferred embodiment, the isolated nucleic acid molecule of the present invention is derived from the aphid *M. persicae* or alternatively, from the Australian sheep blowfly, *L. cuprina*.

30 The ecdysteroid receptor is preferably modulated by one or more of the steroids ecdysone, ponasterone A, or muristerone, or an analogue of an ecdysteroid.

- 19 -

As used herein, the term "analogue of an ecdysteroid" shall be taken to indicate any compound that binds to one or more polypeptide subunits of an ecdysteroid receptor or the heterodimeric holoreceptor comprising same or alternatively or in addition, which binds to the USP polypeptide of a juvenile hormone receptor or alternatively or in addition, which binds to a bioactive derivative or analogue of said polypeptides or holoreceptor. The term "analogue of an ecdysteroid" shall further be taken to indicate any compound that modulates the bioactivity of one or more polypeptide subunits of an ecdysteroid receptor or the heterodimeric holoreceptor comprising same or alternatively or in addition, that modulates the bioactivity of the USP polypeptide of a juvenile hormone receptor or alternatively or in addition, that modulates the bioactivity of modulates the bioactivity of a bioactive derivative or analogue of said polypeptides or holoreceptor.

The present invention is not to be limited in scope to the specific *L. cuprina* and *M. persicae* nucleotide and amino acid sequences set forth in the accompanying Sequence Listing and persons skilled in the art will readily be able to identify additional related sequences from other sources using art-recognised procedures, for example using nucleic acid hybridisation and/or polymerase chain reaction essentially as described by Ausubel *et al.* (1992) and/or McPherson *et al.* (1991) and/or Sambrook *et al.* (1989).

20 Accordingly, the present invention clearly encompasses isolated nucleic acid molecules which encode or are complementary to isolated nucleic acid molecules which encode the subject EcR polypeptide of a steroid receptor or fragments thereof, and/or the subject EcR partner proteins (USP polypeptide) of a steroid receptor and/or the subject USP polypeptide of a juvenile hormone receptor, in addition to derivatives, fragments and analogues thereof which comprise amino acid sequences having at least 40% identity to the amino acid sequences set forth in any one of <400>2, <400>4, <400>6, <400>10, <400>12 or <400>14.

The present invention clearly extends further to isolated nucleic acid molecules which encode or are complementary to isolated nucleic acid molecules which encode the subject EcR polypeptide of a steroid receptor or fragments thereof, and/or the subject EcR partner proteins (USP polypeptide) of a steroid receptor and/or the subject USP polypeptide of a juvenile

- 20 -

hormone receptor, in addition to derivatives, fragments and analogues thereof which comprise amino acid sequences having at least 40% identity to any one or more of the amino acid sequences encoded by the *L. cuprina* or *M. persicae* cDNAs contained in AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568.

5

For the purposes of nomenclature, plasmid pLcEcR contains the cDNA encoding the EcR polypeptide subunit of the *Lucillia cuprina* ecdysone receptor. This plasmid was deposited on 1 July, 1999 with the Australian Government Analytical Laboratories at 1 Suakin Street, Pymble, New South Wales 2073, Australia under the provisions of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and accorded AGAL Accession No. NM99/04566.

For the purposes of nomenclature, plasmid pLcUSP contains the cDNA encoding the EcR partner protein (USP polypeptide) subunit of the *Lucillia cuprina* ecdysone receptor and/or the USP polypeptide subunit of the *L. cuprina* juvenile hormone receptor. This plasmid was deposited on 1 July, 1999 with the Australian Government Analytical Laboratories at 1 Suakin Street, Pymble, New South Wales 2073, Australia under the provisions of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and accorded AGAL Accession No. NM99/04565.

20

For the purposes of nomenclature, plasmid pMpEcR contains the cDNA encoding the EcR polypeptide subunit of the *Myzus persicae* ecdysone receptor. This plasmid was deposited on 1 July, 1999 with the Australian Government Analytical Laboratories at 1 Suakin Street, Pymble, New South Wales 2073, Australia under the provisions of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and accorded AGAL Accession No. NM99/04567.

For the purposes of nomenclature, plasmid pMpUSP contains the cDNA encoding the EcR partner protein (USP polypeptide) subunit of the *Myzus persicae* ecdysone receptor and/or the USP polypeptide subunit of the *M. persicae* juvenile hormone receptor. This plasmid was deposited on 1 July, 1999 with the Australian Government Analytical Laboratories at 1 Suakin

Street, Pymble, New South Wales 2073, Australia under the provisions of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and accorded AGAL Accession No. NM99/04568.

5 The deposits referred to herein will be maintained under the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure. These deposits are provided merely for the purposes of exemplification and are not an admission that a deposit is requird under 35USC §112. A license may be required to make, use or sell the deposited materials or a polypeptide encoded by a cDNA thereof and no 10 such license is hereby granted. It is to be understood however, that the deposits will become publicly available upon the grant of a patent pertaining to the instant disclosure in so far as that patent relates to the deposits referred to herein.

Preferably, the percentage similarity to any one of <400>2, <400>4, <400>6, <400>10, 15 <400>12 or <400>14, or to a polypeptide encoded by a cDNA contained in any one of AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568 is at least about 60%, more preferably at least about 80%, even more preferably at least about 90%.

In determining whether or not two amino acid sequences fall within these percentage limits, 20 those skilled in the art will be aware that it is necessary to conduct a side-by-side comparison or multiple alignment of sequences. In such comparisons or alignments, differences will arise in the positioning of non-identical residues, depending upon the algorithm used to perform the alignment. In the present context, reference to a percentage identity or similarity between two or more amino acid sequences shall be taken to refer to the number of identical and similar 25 residues respectively, between said sequences as determined using any standard algorithm known to those skilled in the art. For example, amino acid sequence identities or similarities may be calculated using the GAP programme and/or aligned using the PILEUP programme of the Computer Genetics Group, Inc., University Research Park, Madison, Wisconsin, United States of America (Devereaux et al, 1984). The GAP programme utilizes the algorithm of 30 Needleman and Wunsch (1970) to maximise the number of identical/similar residues and to minimise the number and/or length of sequence gaps in the alignment. Alternatively or in

addition, wherein more than two amino acid sequences are being compared, the ClustalW programme of Thompson *et al* (1994) is used.

In an alternative embodiment, the isolated nucleic acid molecule of the invention encodes or is complementary to an isolated nucleic acid molecule which encodes a steroid receptor polypeptide or a fragment thereof, or a partner protein (USP) or a fragment thereof, which at least comprises an amino acid sequence which is substantially identical to any one of <400>2, <400>4, <400>6, <400>10, <400>12 or <400>14 or to the amino acid sequence of a polypeptide encoded by any one of the cDNAs contained in AGAL Accession Nos. 10 NM99/04565, NM99/04566, NM99/04567, or NM99/04568.

As used herein, the term "substantially identical" or similar term shall be taken to include any sequence which is at least about 95% identical and preferably at least 99% or 100% identical to a stated nucleotide sequence or amino acid sequence, including any homologue, analogue or derivative of said stated nucleotide sequence or amino acid sequence.

Those skilled in the art will be aware that variants of the nucleotide sequences set forth in any one of <400>1 or <400>3 or <400>5 or <400>7 or <400>8 or <400>9 or <400>11 or <400>13, or variants of the cDNAs contained in any one of the deposited plasmids, which variants encode EcR polypeptides of insect steroid receptors or fragments thereof or EcR partner proteins (USP polypeptides) or fragments thereof, or USP polypeptides of insect juvenile hormone receptors, may be isolated by hybridization under low stringency conditions as exemplified herein.

25 Such variants include any genomic sequences, cDNA sequences mRNA or other isolated nucleic acid molecules derived from the nucleic acid molecules exemplified herein by the Sequence Listing. Additional variants are not excluded.

In a particularly preferred embodiment of the invention, the variant nucleotide sequences encode a fragment of the EcR polypeptide of the insect steroid receptor or a fragment of the EcR partner protein (USP polypeptide) of the insect steroid receptor or a fragment of the USP

- 23 -

polypeptide of the insect juvenile hormone receptor.

Preferred fragments of the subject polypeptides include one or more regions or domains which are involved in the interaction or association between the monomeric polypeptide subunits of a multimeric receptor and/or which are involved in the interaction or association between (i) a cognate steroid or receptor ligand or *cis*-acting DNA sequence; and (ii) said monomeric polypeptide subunits or the receptor *per se.* In a particularly preferred embodiment, the fragments comprise the DNA-binding domain, linker domain (domain D) or a part thereof, or ligand-binding domain (eg. hormone-binding domain) of a steroid receptor polypeptide or juvenile hormone receptor polypeptide or receptor holoenzyme. As exemplified herein, wherein biological activity of the *L. cuprina* ecdysone receptor is required, it is preferably to include at least a ligand-binding region comprising the ligand-binding domain and at least a part of the linker domain of the EcR polypeptide subunit, optionally in association with a ligand-binding region comprising at least the ligand-binding domain and at least a part of the linker domain of the EcR partner protein (USP polypeptide) subunit of said receptor. Additional fragments are not excluded.

Homologues, analogues and derivatives of the nucleotide sequences exemplified herein may be isolated by hybridising same under at least low stringency conditions and preferably under at least medium stringency conditions, to the nucleic acid molecule set forth in any one of <400>1 or <400>3 or <400>5 or <400>7 or <400>8 or <400>9 or <400>11 or <400>13 or to a complementary strand thereof, or to the cDNAs contained in any one or more of the deposited plasmids. More preferably, the isolated nucleic acid molecule according to this aspect of the invention is capable of hybridising under at least high stringency conditions to the nucleic acid molecule set forth in any one of <400>1 or <400>3 or <400>5 or <400>7 or <400>8 or <400>9 or <400>11 or <400>13 or to a complementary strand thereof, or to the cDNAs contained in any one or more of the deposited plasmids.

For the purposes of defining the level of stringency, a low stringency is defined herein as being a hybridisation and/or a wash carried out in 6xSSC buffer, 0.1% (w/v) SDS at 28°C or alternatively, as exemplified herein. Generally, the stringency is increased by reducing the

concentration of SSC buffer, and/or increasing the concentration of SDS and/or increasing the temperature of the hybridisation and/or wash. A medium stringency comprises a hybridisation and/or a wash carried out in 0.2xSSC-2xSSC buffer, 0.1% (w/v) SDS at 42°C to 65°C, while a high stringency comprises a hybridisation and/or a wash carried out in 0.1xSSC-0.2xSSC 5 buffer, 0.1% (w/v) SDS at a temperature of at least 55°C. Conditions for hybridisations and washes are well understood by one normally skilled in the art. For the purposes of further clarification only, reference to the parameters affecting hybridisation between nucleic acid molecules is found in Ausubel et al. (1992), which is herein incorporated by reference.

10 In an even more preferred embodiment of the invention, a hybridising nucleic acid molecule further comprises a sequence of nucleotides which is at least 40% identical to at least 10 contiguous nucleotides, preferably at least 50 contiguous nucleotides and more preferably at least 100 contiguous nucleotides, derived from any one of <400>1 or <400>3 or <400>5 or <400>7 or <400>8 or <400>9 or <400>11 or <400>13 or a complementary strand thereof, or 15 from a cDNA contained in any one or more of the deposited plasmids.

In determining whether or not two nucleotide sequences fall within these percentage limits, those skilled in the art will be aware that it is necessary to conduct a side-by-side comparison or multiple alignment of sequences. In such comparisons or alignments, differences may arise 20 in the positioning of non-identical residues, depending upon the algorithm used to perform the alignment. In the present context, reference to a percentage identity between two or more nucleotide sequences shall be taken to refer to the number of identical residues between said sequences as determined using any standard algorithm known to those skilled in the art. For example, nucleotide sequences may be aligned and their identity calculated using the 25 BESTFIT programme or other appropriate programme of the Computer Genetics Group, Inc., University Research Park, Madison, Wisconsin, United States of America (Devereaux et al, 1984).

In an alternative embodiment, nucleotide sequences encoding EcR polypeptide subunits of 30 insect steroid receptors or fragments thereof and/or EcR partner proteins (USP polypeptides) of insect steroid receptor or fragments thereof, or USP polypeptides of insect juvenile hormone

receptor polypeptides, are amplified in the polymerase chain reaction. According to this embodiment, one or two or more nucleic acid "primer molecules" derived from a nucleotide sequence exemplified herein as <400>1 or <400>3 or <400>5 or <400>7 or <400>8 or <400>9 or <400>11 or <400>13 or a complementary strand thereof, or from a cDNA contained in any one or more of the deposited plasmids, are annealed or hybridized to a nucleic acid "template molecule" which at least comprises a nucleotide sequence encoding a related genetic sequence or a functional part thereof, and nucleic acid molecule copies of the template molecule are amplified enzymatically using a thermostable DNA polymerase enzyme, such as Tagl polymerase or Pfu polymerase, amongst others.

10

More particularly, one of the primer molecules comprises contiguous nucleotides derived from any one of <400>1 or <400>3 or <400>5 or <400>7 or <400>8 or <400>9 or <400>11 or <400>13, or alternatively, from a cDNA contained in any one or more of the deposited plasmids; and another of said primers comprises contiguous nucleotides complementary to <400>1 or <400>3 or <400>5 or <400>7 or <400>8 or <400>9 or <400>11 or <400>13, or alternatively, from a cDNA contained in any one or more of the deposited plasmids, subject to the proviso that the first and second primers are not complementary to each other.

In a preferred embodiment, each nucleic acid primer molecule is at least 10 nucleotides in length, more preferably at least 20 nucleotides in length, even more preferably at least 30 nucleotides in length, still more preferably at least 40 nucleotides in length and even still more preferably at least 50 nucleotides in length.

Furthermore, the nucleic acid primer molecules consists of a combination of any of the nucleotides adenine, cytidine, guanine, thymidine, or inosine, or functional analogues or derivatives thereof which are at least capable of being incorporated into a polynucleotide molecule without having an inhibitory effect on the hybridisation of said primer to the template molecule in the environment in which it is used.

30 Furthermore, one or both of the nucleic acid primer molecules may be contained in an aqueous mixture of other nucleic acid primer molecules, for example a mixture of degenerate

primer sequences which vary from each other by one or more nucleotide substitutions or deletions. Alternatively, one or both of the nucleic acid primer molecules may be in a substantially pure form.

In a particularly preferred embodiment exemplified herein, two primer nucleotide sequences are used to amplify related sequences, said primers comprising the nucleotide sequences as set forth in any one of <400>15 to <400>20 inclusive. Even more preferably, the primers are used in the combination of (i)<400>15 and <400>16; or (ii) <400>17 and <400>18; or (iii) <400>19 and <400>20.

10

The nucleic acid template molecule may be in a recombinant form, in a virus particle, insect cell, bacteriophage particle, yeast cell, animal cell, or a plant cell. Preferably, the nucleic acid template molecule is derived from an insect species.

Those skilled in the art will be aware that there are many known variations of the basic polymerase chain reaction procedure. Such variations are discussed, for example, in McPherson et al (1991). The present invention extends to the use of all such variations in the isolation of variant insect steroid receptor-encoding genes or fragments thereof, and/or variant partner protein-encoding genes or fragments thereof to those exemplified herein.

20

The isolated nucleic acid molecule of the present invention, including those sequences exemplified herein and any variants thereof, may be cloned into a plasmid or bacteriophage molecule, for example to facilitate the preparation of primer molecules or hybridisation probes or for the production of recombinant gene products. Methods for the production of such recombinant plasmids, cosmids, bacteriophage molecules or other recombinant molecules are well-known to those of ordinary skill in the art and can be accomplished without undue experimentation. Accordingly, the invention further extends to any recombinant plasmid, bacteriophage, cosmid or other recombinant molecule comprising the nucleotide sequence set forth in any one of <400>1 or <400>3 or <400>5 or <400>7 or <400>8 or <400>9 or <400>11 or <400>13 or <400>15 to <400>20, or a complementary sequence, homologue, analogue or derivative thereof, or a cDNA contained in any one or more of the deposited plasmids.

- 27 -

The nucleic acid molecule of the present invention is also useful for developing genetic constructs which comprise and preferably, express, the EcR polypeptide subunit of the insect steroid receptor and/or the EcR partner protein (USP polypeptide) of the steroid receptor and/or the USP polypeptide of the juvenile hormone receptor, thereby providing for the production of the recombinant polypeptides in isolated cells or transformed tissues.

Accordingly, a further aspect of the present invention provides a genetic construct comprising the subject isolated nucleic acid molecule encoding the insect steroid receptor polypeptide or a juvenile hormone receptor polypeptide, operably linked to a promoter sequence. Preferably, the subject nucleic acid molecule is in an expressible format, such that it is possible to produce a recombinant polypeptide therefrom.

Reference herein to a "promoter" is to be taken in its broadest context and includes the transcriptional regulatory sequences of a classical genomic gene, including the TATA box which is required for accurate transcription initiation in a eukaryotic cell, with or without a CCAAT box sequence or alternatively, the Pribnow box required for accurate expression in prokaryotic cells.

Promoters may be cell, tissue, organ or system specific, or may be non-specific. Using specific promoters, the expression of a bioactive agent or other polypeptide encoded by a structural gene to which the promoter is operably connected may be targeted to a desired cellular site. For example, in transgenic animals such as sheep, it can be envisaged that cells of the transgenic animal may contain a gene encoding a steroid receptor, preferably a steroid receptor linked to an epidermal specific promoter and a separate gene encoding, for example, epidermal growth factor (EGF) which is functionally linked to one or more insect hormone response elements and may or may not also be linked to epidermal specific promoter elements. On administration of the appropriate insect steroid hormone to the transgenic animal, the activated complex between the insect steroid receptor and insect steroid may bind to the one or more insect steroid hormone response element thereby inducing EGF production solely in epidermal cells which may give rise to deflecting. It is to be understood that this aspect of the invention is independent of the degree of thermostability of the insect steroid

receptor The same principal applies to expression of any bioactive molecule or reporter molecule in a specific cell type which is regulated by a transactivating complex between a steroid receptor complex and an appropriate insect steroid.

5 In the present context, the term "promoter" is also used to describe a synthetic or fusion molecule, or derivative which confers, activates or enhances expression in a cell in response to an external stimulus. Accordingly, the promoter may include further regulatory elements (i.e. upstream activating sequences, enhancers and silencers) which alter gene expression in response to developmental and/or external stimuli, or in a tissue-specific manner. Preferred promoters may contain copies of one or more specific regulatory elements, in particular steroid responsive elements (SREs) or hormone-responsive elements (HREs), to further enhance expression and/or to alter the spatial expression and/or temporal expression pattern.

Reference herein to the term "steroid response element" shall be taken to refers to one or more *cis*-acting nucleotide sequences present in a naturally-occurring or synthetic or recombinant gene the expression of which is regulated by an insect steroid, such as an ecdysteroid, for example ecdysone or ponasterone A, wherein said regulation of expression results from an direct or indirect interaction between a steroid receptor and said *cis*-acting nucleotide sequence response element. Exemplary insect steroid hormone response elements include the ecdysone response element hsp27 (EcRE) and any other nucleotide sequence which is capable of binding ecdysteroid receptors or polypeptide subunits thereof or fragments or analogies thereof (such as associated with E75, E74 or other *Drosophila* early genes), as described for example by Riddihough and Pelham (1987).

For example, an SRE or a plurality of such elements may be operably linked to a promoter such as the polyhedron promoter, p10 promoter, MMTV promoter or SV40 promoter, to make transcription of a structural gene to which said promoter is operably connected responsive to the presence of a steroid bound to the insect receptor (which may act as a transcription factor). One or more insect SREs may be located within a promoter, and may replace sequences within a selected promoter which confer responsiveness to hormones or other agents which regulate promoter activity. Where response elements are different they may lead to

- 29 -

preferential binding of different insect steroids or analogues thereof such that a promoter may be differentially regulated.

Particularly preferred SREs according to this embodiment include, but are not limited to, the hsp27 ecdysone response element described by Riddihough and Pelham (1987) or the 13 base-pair palindromic core contained therein.

A promoter is usually, but not necessarily, positioned upstream or 5', of a structural gene, the expression of which it regulates. Furthermore, the regulatory elements comprising a promoter are usually positioned within 2 kb of the start site of transcription of the gene.

Placing a gene or isolated nucleic acid molecule operably under the control of a promoter sequence means positioning said gene or isolated nucleic acid molecule such that its expression is controlled by the promoter sequence. Promoters are generally positioned 5' (upstream) to the genes that they control. In the construction of heterologous promoter/structural gene combinations it is generally preferred to position the promoter at a distance from the gene transcription start site that is approximately the same as the distance between that promoter and the gene it controls in its natural setting, i.e., the gene from which the promoter is derived. As is known in the art, some variation in this distance can be accommodated without loss of promoter function. Similarly, the preferred positioning of a regulatory sequence element with respect to a heterologous gene to be placed under its control is defined by the positioning of the element in its natural setting, i.e., the genes from which it is derived. Again, as is known in the art, some variation in this distance can also occur.

25

Those skilled in the art will recognise that the choice of promoter will depend upon the nature of the cell being transformed and when expression is required. Furthermore, it is well-known in the art that the promoter sequence used in the expression vector will also vary depending upon the level of expression required and whether expression is intended to be constitutive or regulated.

For expression in eukaryotic cells, the genetic construct generally comprises, in addition to the nucleic acid molecule of the invention, a promoter and optionally other regulatory sequences designed to facilitate expression of said nucleic acid molecule. The promoter may be derived from a genomic clone which normally encodes the expressed protein or alternatively, it may be a heterologous promoter derived from another genetic source. Promoter sequences suitable for expression of genes in eukaryotic cells are well-known in the art.

Suitable promoters for use in eukaryotic expression vectors include those capable of regulating expression in mammalian cells, insect cells such as Sf9 or Sf21. (Spodoptera frugiperda) cells, 10 yeast cells and plant cells. Preferred promoters for expression in eukaryotic cells include the p10 promoter, MMTV promoter, polyhedron promoter, the SV40 early promoter and the cytomegalovirus (CMV- IE) promoter, promoters derived from immunoglobulin-producing cells (see, United States Patent No 4,663,281), polyoma virus promoters, and the LTR from various retroviruses (such as murine leukemia virus, murine or Rous sarcoma virus and HIV), amongst others (See, Enhancers and Eukaryotic Gene Expression, Cold Spring Harbor Press, New York, 1983, which is incorporated herein by reference). Examples of other expression control sequences are enhancers or promoters derived from viruses, such as SV40, Adenovirus, Bovine Papilloma Virus, and the like.

Wherein the expression vector is intended for the production of recombinant protein, the promoter is further selected such that it is capable of regulating expression in a cell which is capable of performing any post-translational modification to the polypeptide which may be required for the subject recombinant polypeptide to be functional, such as N-linked glycosylation. Cells suitable for such purposes may be readily determined by those skilled in the art. By way of exemplification, Chinese hamster ovary (CHO) cells may be employed to carry out the N-terminal glycosylation and signal sequence cleavage of a recombinant polypeptide produced therein. Alternatively, a baculovirus expression vector such as the pFastBac vector supplied by GibcoBRL may be used to express recombinant polypeptides in Sf9 (Spodoptera frugiperda) cells, following standard protocols.

30

Numerous expression vectors suitable for the present purpose have been described and are

P:\OPER\MRO\ECDYSONE.CIP - 1/7/99

- 31 -

readily available. The expression vector may be based upon the pcDNA3 vector distributed by Medos Company Pty Ltd, Victoria, Australia, which comprises the CMV promoter and BGH terminator sequences for regulating expression of the recombinant polypeptide of the invention in a eukaryotic cell, when isolated nucleic acid sequences encoding same are inserted, in the sense orientation relative to the CMV promoter, into the multiple cloning site of said vector. Alternatively, the SG5 expression vector of Greene et al. (1988), supplied by Stratagene, or the pQE series of vectors supplied by Qiagen are particularly useful for such purposes, as exemplified herein.

- 10 Examples of eukaryotic cells contemplated herein to be suitable for expression include mammalian, yeast, insect, plant cells or cell lines such as COS, VERO, HeLa, mouse C127, Chinese hamster ovary (CHO), WI-38, baby hamster kidney (BHK), MDCK, sf21 (insect) or Sf9 (insect) cell lines. Such cell lines are readily available to those skilled in the art.
- The prerequisite for expression in prokaryotic cells such as Escherichia coli is the use of a strong promoter with an effective ribosome binding site. Typical promoters suitable for expression in bacterial cells such as E. coli include, but are not limited to, the lacz promoter, temperature-sensitive λ_c or λ_c promoters, T7 promoter or the IPTG-inducible tac promoter. A number of other vector systems for expressing the nucleic acid molecule of the invention in 20 E.coli are well-known in the art and are described for example in Ausubel et al (1992).

Numerous vectors having suitable promoter sequences for expression in bacteria have been described, such as for example, pKC30 (λ_L:Shimatake and Rosenberg, 1981), pKK173-3 (*tac*: Amann and Brosius, 1985), pET-3 (T7: Studier and Moffat, 1986) or the pQE series of expression vectors (Qiagen, CA), amongst others.

Suitable prokaryotic cells include corynebacterium, salmonella, *Escherichia coli, Bacillus* sp. and *Pseudomonas* sp. amongst others. Bacterial strains which are suitable for the present purpose are well-known in the relevant art (Ausubel et al, 1992).

The genetic constructs described herein may further comprise genetic sequences

- 32 --

corresponding to a bacterial origin of replication and/or a selectable marker gene such as an antibiotic-resistance gene, suitable for the maintenance and replication of said genetic construct in a prokaryotic or eukaryotic cell, tissue or organism. Such sequences are well-known in the art,

5

Selectable marker genes include genes which when expressed are capable of conferring resistance on a cell to a compound which would, absent expression of said selectable marker gene, prevent or slow cell proliferation or result in cell death. Preferred selectable marker genes contemplated herein include, but are not limited to antibiotic-resistance genes such as those conferring resistance to ampicillin, Claforan, gentamycin, G-418, hygromycin, rifampicin, kanamycin, neomycin, spectinomycin, tetracycline or a derivative or related compound thereof or any other compound which may be toxic to a cell.

The origin of replication or a selectable marker gene will be spatially-separated from those genetic sequences which encode the recombinant receptor polypeptide or fusion polypeptide comprising same.

Preferably, the genetic constructs of the invention, including any expression vectors, are capable of introduction into, and expression in, an *in vitro* cell culture, or for introduction into, with or without integration into the genome of a cultured cell, cell line and/or transgenic animal.

In a particularly preferred embodiment, the expression vector is selected from the group consisting of: pLcEcR (AGAL Accession No. NM99/04566); pLcUSP (AGAL Accession No. NM99/04565); pMpEcR (AGAL Accession No. NM99/04567); and pMpUSP (AGAL Accession No. NM99/04568).

A further aspect of the invention provides a cell comprising the subject isolated nucleic acid molecule which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide.

30

As used herein, the word "cell" shall be taken to refer to a single cell, or a cell lysate, or a

tissue, organ or whole organism comprising same, including a tissue, organ or whole organism comprising a clonal group of cells or a heterogenous mixture of cell types, which may be a prokaryotic or eukaryotic cell as described *supra*.

5 in a preferred embodiment, the cell of the present invention expresses the isolated or recombinant polypeptide encoded by the nucleic acid molecule.

In a preferred embodiment, the cell expresses a steroid receptor polypeptide or a fragment thereof which receptor is capable of binding to an insect steroid or analogue thereof or a candidate insecticidally active agent to form an activated complex, and comprises a nucleic acid sequence encoding a bioactive molecule or a reporter molecule operably linked to one or more insect steroid response elements which on binding of the said activated complex promotes transcription of the nucleic acid sequence, wherein said cell on exposure to insect steroid or an analogue thereof, regulates expression of said bioactive molecule or allows detection of said reporter molecule.

To produce the cells of the invention, host cells are transfected or co-transfected or transformed with nucleotide sequences containing the DNA segments of interest (for example, the insect steroid receptor gene, the recombinant steroid response elements, or both) by well-known methods, which vary depending on the type of cellular host. For example, calcium chloride transfection is commonly utilized for prokaryotic cells, whereas lipofection or calcium phosphate treatment are often used for other cellular hosts. See, generally, Sambrook et al, (1989); Ausubel et al, (1992); and Potrykus (1990). Other transformation techniques include electroporation, DEAE-dextran, microprojectile bombardment, lipofection, microinjection, and others.

As used herein, the term "transformed cell" is meant to also include the progeny of a transformed cell.

30 In a further aspect of this invention, there is provided an animal (such as a mammal or insect), microorganism, plant or aquatic organism, containing one or more cells as mentioned above.

20

P:\OPER\MEO\ECDYSONE.CIP - 1/7/99

Reference to plants, microorganisms and aquatic organisms includes any such organisms.

In this embodiment of the invention, it is to be appreciated that administration of an insect steroid or an analogue thereof to an organism will induce expression of the desired bioactive molecule, such as a polypeptide, with attendant advantages. For example, an induced protein may have a therapeutic effect ameliorating a disease state or preventing susceptibility to disease or may modify in some way the phenotype of an organism to produce a desired effect. In humans, for example, cell transplants (such as liver cells) may under the action of insect steroids, produce desirable hormones such as insulin, growth hormone, growth factors and the like.

A further aspect of the invention provides a recombinant or isolated polypeptide comprising a steroid receptor polypeptide or juvenile hormone receptor polypeptide derived from an insect or a bioactive derivative or analogue thereof, wherein said polypeptide:

- is selected from the list comprising EcR polypeptide of a steroid receptor, the partner protein (USP polypeptide) of a steroid receptor and the USP polypeptide of a juvenile hormone receptor; and
 - (ii) comprises an amino acid sequence that is at least 40% identical to any one of the amino acid sequences set forth in <400>2, <400>4, <400>6, <400>10, <400>12 or <400>14;

wherein said polypeptide is substantially free of naturally-associated insect cell components.

In an alternative embodiment, the recombinant or isolated polypeptide comprising a steroid receptor polypeptide or juvenile hormone receptor polypeptide derived from an insect or a bioactive derivative or analogue thereof, wherein said polypeptide:

- (i) is selected from the list comprising EcR polypeptide of a steroid receptor, the partner protein (USP polypeptide) of a steroid receptor and the USP polypeptide of a juvenile hormone receptor; and
- (ii) comprises an amino acid sequence that is at least 40% identical to an amino
 30 acid sequence encoded by the cDNA present in any one of the plasmids deposited
 under AGAL Accession No. NM99/04565, NM99/04566, NM99/04567, or NM99/04568;

- 35 -

wherein said polypeptide is substantially free of naturally-associated insect cell components.

Reference herein to "substantially free of naturally associated insect cell components" refers to at least 80% purity, preferably more than 90% purity, and more preferably more than 95% 5 purity. Normally, purity is measured on a polyacrylamide gel with homogeneity determined by staining of protein bonds. Alternatively, high resolution may be necessary using HPLC or similar means. For most purposes, a simple chromatography column or polyacrylamide gel may be used to determine purity. A protein which is chemically synthesized or synthesized in a cell system different from an insect cell from which it naturally originates would be free of 10 naturally-associated insect cell components.

The present invention clearly provides for the isolation of EcR polypeptide subunits and EcR partner protein (USP polypeptide) subunits of ecdysteroid receptors and USP polypeptides of juvenile hormone receptors, from various organisms of the class Insecta, as described supra, 15 in addition to protozoa and helminth sources.

Insect steroid receptors are characterized by functional ligand-binding domains, and DNAbinding domains, both of which interact to effect a change in the regulatory state of a gene operably linked to the DNA-binding site of the holoreceptor or a polypeptide or polypeptide 20 fragment thereof. Thus, insect steroid receptors seem to be ligand-responsive transcription factors. Additionally, insect steroid receptors generally contain a DNA-binding domain (Domain C), and a ligand-binding domain (Domain E), separated and flanked by additional domains as identified by Krust et al (1986). The C domain preferably comprises a zinc-finger DNA-binding domain which is usually hydrophilic, having high cysteine, lysine and arginine 25 content. The E domain preferably comprises hydrophobic amino acid residues and is further characterized by regions E1, E2 and E3. The ligand-binding domain of the members of the insect steroid receptor superfamily is typically carboxyl-proximal, relative to a DNA-binding domain (Evans, 1988). The entire ligand-binding domain is typically between about 200 and 250 amino acids but is potentially shorter. This domain has the subregions of high homology, 30 designated the E1, E2 and E3 regions - which may be collectively referred to as the "E region". Amino acid residues proximal to the C domain comprise a region initially defined as separate

P:\OPER\MRO\ECDYSONE,CIP - 1/7/99

- 36 -

A and B domains. Region D separates the more conserved domains C and E. Region D typically has a hydrophilic region whose predicted secondary structure is rich in turns and coils. The F region is carboxy promixal to the E region (see, Krust *et al, supra*).

5 The receptor polypeptides of the present invention exhibit at least a ligand-binding domain, as characterized by sequence homology to regions E1, E2 and E3. The ligand-binding domains of the present invention are typically characterized by having significant homology in sequence and structure to these three regions. Fragments of insect steroid receptors and partner proteins capable of binding insect steroids, and candidate insecticidally active compounds comprise an E-region or a sufficient portion of the E-region to allow binding.

Preferably, the recombinant or isolated EcR polypeptide subunit of the insect steroid receptor or EcR partner protein (USP polypeptide) subunit of the steroid receptor or USP polypeptide of the juvenile hormone receptor as described herein is thermostable.

By "thermostable" is meant that a stated integer does not exhibit reduced activity at bacterial, plant or animal physiological temperatures above about 28°C or above about 30°C. The thermostability of insect steroid hormone receptors also refers to the capacity of such receptors to bind to ligand-binding domains or regions and/or to transactivate genes linked to 20 insect steroid hormone response elements at bacterial, plant or animal physiological temperatures above about 28°C or above about 30°C.

The present invention clearly extends to variants of said polypeptides, as described *supra*. The polypeptide may be substantially free of naturally associated insect cell components, or may be in combination with a partner protein which associates with the insect steroid receptor so as to confer enhanced affinity for insect steroid response elements, enhanced affinity for insect steroids or analogues thereof. For Example, the amino acid sequences exemplified herein may be varied by the deletion, substitution or insertion of one or more amino acids.

30 In one embodiment, amino acids of a polypeptide exemplified herein may be replaced by other amino acids having similar properties, for example hydrophobicity, hydrophilicity, hydrophobic

P:\OPER\MRO\ECDYSONE,CIP - 1/7/99

- 37 -

moment, charge or antigenicity, and so on.

Substitutions encompass amino acid alterations in which an amino acid of the base polypeptide is replaced with a different naturally-occurring or a non-conventional amino acid 5 residue. Such substitutions may be classified as "conservative", in which case an amino acid residue contained in the base polypeptide is replaced with another naturally-occurring amino acid of similar character, for example Gly↔Ala, Val↔Ile↔Leu, Asp↔Glu, Lys↔Arg, Asn↔Gln or Phe↔Trp↔Tyr.

10 Substitutions encompassed by the present invention may also be "non-conservative", in which an amino acid residue which is present in the base polypeptide is substituted with an amino acid having different properties, such as a naturally-occurring amino acid from a different group (eg. substituted a charged or hydrophobic amino acid with alanine), or alternatively, in which a naturally-occurring amino acid is substituted with a non-conventional amino acid.

Those skilled in the art will be aware that several means are available for producing variants of the exemplified EcR polypeptide subunit of the insect steroid receptor or EcR partner protein (USP polypeptide) subunit of the steroid receptor or USP polypeptide of the juvenile hormone receptor, when provided with the nucleotide sequence of the nucleic acid molecule which 20 encodes said polypeptide, for example site-directed mutagenesis of DNA and polymerase chain reaction utilising mutagenised oligonucleotide primers, amongst others.

Such polypeptide variants which are capable of binding insect steroids clearly form part of the present invention. Assays to determine such binding may be carried out according to 25 procedures well known in the art.

One such variant polypeptide encompassed by the present invention comprises an "in-frame" fusion polypeptide between different regions of different insect receptor polypeptides. As exemplified herein, the present inventors have discovered that, by producing synthetic genes 30 in which various domains of a base insect steroid receptor-encoding nucleotide sequence derived from a first source are interchanged or substituted with similar sequences derived from

25

30

P:\OPER\MRO\ECDYSONE.CIP - 1/7/99

a second source (referred to as "domain swapping"), it is possible to modify the bioactivity of the insect steroid receptor encoded therefor. For example, the biological activity of the EcR polypeptide of the *L. cuprina* or *M. persicae* ecdysone receptor exemplified herein may be modulated by replacing portions of its C-terminal or N-terminal sequences with the equivalent domains from the EcR polypeptide of the *D. melanogaster* ecdysone receptor or alternatively, by swapping regions of the EcR polypeptides of the *L. cuprina* and *M. persicae* ecdysone receptors *per se*.

As a further refinement, such changes in biological function can similarly be effected by making specific changes (e.g. addition, substitution or deletion) to only those amino-acids within each domain that are critical for determining the relevant catalytic function (eg. ligand-binding activity, DNA binding site affinity, etc), such as by site-directed mutagenesis.

According to this embodiemtn, there is provided a synthetic EcR polypeptide subunit of a steroid receptor, and/or a synthetic EcR partner protein (USP polypeptide) subunit of a steroid receptor, and/or a synthetic USP polypeptide of a juvenile hormone receptor, or an analogue or derive of said synthetic polypeptides, wherein said synthetic polypeptides comprise an amino acid sequence which has the following properties:

- (i) it differs in amino acid sequence or exhibits different biological properties to a naturally-occurring EcR polypeptide subunit of a steroid receptor, and/or a naturally-occurring EcR partner protein (USP polypeptide) subunit of a steroid receptor, and/or a naturally-occurring USP polypeptide of a juvenile hormone receptor;
 - (ii) it comprises a first sequence of amino acids which are at least about 40% identical to a part of any one of <400>2, <400>4, <400>6, <400>10, <400>12 or <400>14, or at least about 40% identical to a part of an amin oa cid sequence encoded by any one of the deposited plasmids, linked covalently to a second sequence of amino acids derived from an EcR polypeptide subunit of a steroid receptor, EcR partner protein (USP polypeptide) subunit of a steroid receptor, or USP polypeptide of a juvenile hormone receptor, wherein said first and second sequences are derived from different genomic sources.

- 39 -

Preferably, the first sequence of amino acids is derived from the EcR polypeptide subunit of a steroid receptor, more preferably from the EcR polypeptide of the L. cuprina or M. persicae ecdysone receptor, and even more preferably from the the EcR polypeptide of the L. cuprina ecdysone receptor.

5

In one embodiment, the synthetic EcR polypeptide subunit of a steroid receptor, and/or a synthetic EcR partner protein (USP polypeptide) subunit of a steroid receptor, and/or a synthetic USP polypeptide of a juvenile hormone receptor comprises a fusion polypeptide in which the ligand-binding regions of any one of <400>2, <400>4, <400>6; <400>10, <400>12 10 or <400>14 are replaced, in-frame, by the ligand-binding region of a different receptor polypeptide...

In a particularly preferred embodiment, 5'-end of the open reading frame of a first nucleotide sequence, encoding the N-terminal portion of the EcR polypeptide of a first ecdysteroid 15 receptor to the end of the DNA-binding domain of said polypeptide, is fused in-frame, to the 3'-end of the open reading frame of a second nucleotide sequence, encoding the C-terminal portion of the EcR polypeptide of a second ecdysteroid receptor, from the D domain and hormone-binding domain to the carboxyl terminus.

20 Accordingly, the present invention extends to any variants of the insect receptor polypeptides referred to herein and genetic sequences encoding same, wherein said variants are derived from a receptor polypeptide as described herein and exhibit demonstrable ligand-binding activity, and either comprises an amino acid sequence which differs from a naturally-occurring receptor polypeptide, or exhibit biological activity.

25

As with other aspects of the invention, the variants described herein may be produced as recombinant polypeptides or in transgenic organisms, once the subject synthetic genes are introduced into a suitable host cell and expressed therein.

30 In an alternative embodiment, the recombinant receptor polypeptide of the invention is produced as an "in-frame" fusion polypeptide with a second polypeptide, for example a

- 40 -

detectable reporter polypeptide such as β -galactosidase, β -glucuronidase, luciferase or other enzyme, or a FLAG peptide, hapten peptide such as a poly-lysine or poly-histidine or other polypeptide molecule.

5 By "in-frame" means that a nucleotide sequence which encodes a first polypeptide is placed (i.e. cloned or ligated) in the same open reading frame adjacent to a nucleotide sequence which encodes a second polypeptide with no intervening stop codons there between, such that when the ligated nucleic acid molecule is expressed, a single fusion polypeptide is produced which comprises a sequence of amino acids corresponding to the summation of the individual amino acid sequences of the first and second polypeptides.

In order to produce a fusion polypeptide, the nucleic acid molecule which encodes the polypeptide of the invention, or an analogue or derivative thereof, is cloned adjacent to a second nucleic acid molecule encoding the second polypeptide, optionally separated by a spacer nucleic acid molecule which encodes one or more amino acids (eg: poly-lysine or poly histidine, amongst others), such that the first coding region and the second coding region are in the same open reading frame, with no intervening stop codons between the two coding regions. When translated, the polypeptide thus produced comprises a fusion between the polypeptide products of the first and second coding regions. Wherein a spacer nucleic acid molecule is utilised in the genetic construct, it may be desirable for said spacer to at least encode an amino acid sequence which is cleavable to assist in separation of the fused polypeptide products of the first and second coding regions, for example a thrombin cleavage site.

25 A genetic construct which encodes a fusion polypeptide further comprises at least one start codon and one stop codon, capable of being recognised by the cell's translational machinery in which expression is intended.

Preferably, a genetic construct which encodes a fusion polypeptide may be further modified to include a genetic sequence which encodes a targeting signal placed in-frame with the coding region of the nucleotide sequence encoding the fusion polypeptide, to target the

- 41 -

expressed recombinant polypeptide to the extracellular matrix or other cell compartment. More preferably, the genetic sequence encoding targeting signal is placed in-frame at the 5'-terminus or the 3'-terminus, but most preferably at the 5'-terminus, of the coding region of the nucleotide sequence which encodes the fusion polypeptide.

5

Methods for the production of a fusion polypeptide are well-known to those skilled in the art.

The recombinant EcR polypeptide subunit of the insect steroid receptor or EcR partner protein (USP polypeptide) subunit of the steroid receptor or USP polypeptide of the juvenile hormone receptor may be purified by standard techniques, such as column chromatography (using various matrices which interact with the protein products, such as ion exchange matrices, hydrophobic matrices and the like), affinity chromatography utilizing antibodies specific for the protein or other ligands such as dyes or insect steroids which bind to the protein.

15 Wherein the recombinant polypeptide is expressed as a fusion polypeptide, it is also possible to purify the fusion polypeptide based upon its properties (eg size, solubility, charge etc). Alternatively, the fusion polypeptide may be purified based upon the properties of the non-receptor moiety of said fusion polypeptide, for example substrate affinity. Once purified, the fusion polypeptide may be cleaved to release the intact polypeptide of the invention.

20

Alternatively, proteins may be synthesized by standard protein synthetic techniques as are well known in the art.

In a preferred embodiment, the recombinant or isolated polypeptides of the invention are provided as a precipitate or crystallized by standard techniques, preferably for X-ray crystal structure determination.

The three-dimensional structure of the polypeptide of the invention or a holoreceptor comprising same or a fragment of said polypeptide or holoreceptor is particularly useful for identifying candidate insecticidal agents which mimic ligands that bind to said three-dimensional structure and/or modulate the ability of insect steroids to bind thereto and activate

P:\OPER\MRO\ECDYSONE.CIP - 1/7/99

- 42 -

the receptor (see, for example, Von Itzstein et al., 1993; and Bugg et al., 1993).

According to this embodiment, the EcR polypeptides of the invention or ligand binding domains thereof, or their complexes with EcR partner proteins or ligand binding domains thereof, which confer enhanced affinity for insect steroid response elements or partner proteins (USP polypeptides) or ligands, are particularly useful to model the three-dimensional structure of the receptor ligand-binding region. In this manner, insecticidal compounds may be produced which bind to, or otherwise interact with, the ligand-binding region of the receptor and/or preferably interfere with ligand binding. In the same way, compounds may be developed which have a potentiated interaction with the insect steroid receptor over and above that of the physiological insect steroid which binds to the receptor.

Accordingly, a still further aspect of the invention provides a method of identifying a candidate insecticidally-active agent comprising the steps of:

- a) expressing a USP polypeptide of a juvenile hormone receptor or a fragment thereof which includes the ligand-binding region, optionally in association with an EcR polypeptide of a steroid receptor or ligand binding domain thereof, and optionally in association with an insect steroid or analogue thereof, so as to form a complex;
 - b) purifying or precipitating the complex:
- c) determining the three-dimensional structure of the ligand binding domain of the complex; and
 - d) identifying compounds which bind to or associate with the three-dimensional structure of the ligand binding domain, wherein said compounds represent candidate insecticidally-active agents.

Standard procedures are used to determine the three dimensional structure of the receptor polypeptides of the invention, for example using X-ray crystallography and/or nuclear magnetic resonance analysis (see, for example, Bugg et al., 1993; Von Itstein et al., 1993).

30 Insecticidally-active agents contemplated herein include synthetic chemicals that mimic one or more ligands of the holoreceptor or its polypeptide subunit, or the ligand-binding region of - 43 -

said holoreceptor or subunit, thereby modulating binding of steroids to said holoreceptor or subunit. Preferred insecticidally-active agents include bisacylhydrazines, iridoid glycosides or other non-steroidal modulators of ecdysteroid receptors or insect juvenile hormone receptors. Additionally, because the EcR partner protein (USP polypeptide) subunits of insect steroid receptors, and the USP polypeptides of insect juvenile hormone receptors, bind insect juvenile hormones, a sesquiterpenoid group of ligands that regulate developmental transitions in insects (see Jones and Sharp,1997), compounds which interfere with the binding of juvenile hormone are also candidate insecticides.

- 10 A further aspect of the present invention provides a method of identifying a modulator of insect steroid receptor-mediated gene expression or insect juvenile hormone receptor-mediated gene expression comprising:
 - (i) assaying the expression of a reporter gene in the presence of a recombinant or isolated insect steroid receptor polypeptide or a juvenile hormone receptor polypeptide of the invention and a potential modulator; and
 - (ii) assaying the expression of a reporter gene in the presence of a recombinant or isolated insect steroid receptor polypeptide or a juvenile hormone receptor polypeptide of the invention and without said potential modulator; and
 - (ii) comparing expression of the reporter gene in the presence of the potential modulator to the expression of a reporter gene in the absence of the potential modulator,

wherein said reporter gene is placed operably under the control of a steroid response element (SRE) to which said insect steroid receptor binds or a promoter sequence comprising said SRE.

25

15

20

In the present context, a "modulator" is a compound or molecule that agonises or antagonises the binding properties and/or biological activity of a receptor polypeptide or holoreceptor. Preferred modulators according to this embodiment include those synthetic compounds that are suitable for use as insecticidally-active agents described *supra*.

30

The reporter gene may be any gene, the expression of which may be monitored or assayed

- 44 -

readily. Preferably, the reporter gene is a structural gene that encodes a peptide, polypeptide or enzyme that is assayed readily by enzymic or immunological means, for example the βgalactosidase, β-glucuronidase, luciferase or chloramphenicol acetyltransferase (CAT) genes. Alternatively, the reporter gene may be a gene which encodes an immunologically-detectable 5 protein, for example a FLAG peptide, poly-lysine peptide or poly-histidine peptide.

Standard methods are used to assay the expression of the reporter gene.

This embodiment of the invention may be applied directly to the identification of potential 10 insecticidally-active compounds or alternatively, modified for such purposes by assaying for the binding (direct or indirect) of the recombinant or isolated insect steroid receptor polypeptide or a juvenile hormone receptor polypeptide of the invention to a steroid response element (SRE), rather than by assaying for reporter gene expression. According to this alternative embodiment, the binding assayed in the presence or absence of a potential 15 insecticidally-active compound is compared, wherein a difference in the level of binding indicates that the candidate compound possesses potential insecticidal activity.

In addition, substances may be screened for insecticidal activity by assessing their ability to bind, in vivo or in vitro, to the intact ecdysone receptor or alternatively, the ligand-binding 20 regions of the EcR polypeptide subunit of the ecdysone receptor (eg. <400>2 and/or <400>6 and/or <400>10) and/or the EcR partner protein (USP polypeptide) of the ecdysone receptor (eg. <400>4 and/or <400>12 and/or <400>14). Competition assays involving the native insect steroid may be employed to assess insecticidal activity.

25 The performance of this embodiment may, for example, involve binding the insect steroid receptor polypeptide to a support such as a plurality of polymeric pins, whereafter the polypeptide resident on the plurality of pins is brought into contact with candidate insecticidal molecules for screening. The molecules being screened may be isotopically labelled so as to permit ready detection of binding. Alternatively, reporter molecules may be utilized which bind 30 to the insect steroid receptor candidate molecule complex. Alternatively, compounds for screening may be bound to a solid support, such as a plurality of pins which are then reacted

- 45 -

with the thermostable insect steroid receptor or complex with a partner protein. Binding may, for example, be determined again by isotopic-labelling of the receptor, or by antibody detection or use of another reporting agent.

5 In an alternative embodiment, insecticidally-active agent are identified using rational drug design, by expressing a USP polypeptide of a juvenile hormone receptor or a fragment thereof which includes the ligand-binding region, optionally in association with an EcR polypeptide of a steroid receptor or ligand binding domain thereof, and optionally in association with an insect steroid or analogue thereof, so as to form a complex, determining the three-dimensional 10 structure of the ligand binding domain of the complex, and identifying compounds which bind to or associate with the three-dimensional structure of the ligand binding domain, wherein said compounds represent candidate insecticidally-active agents.

The methods described herein for identifying modulators of gene expression and insecticidal 15 compounds, may be performed using prokaryotic or eukaryotic cells, cell lysates or aqueous solutions.

A further aspect of this invention accordingly relates to synthetic compounds derived from the three dimensional structure of EcR polypeptides and/or EcR partner protein (USP polypeptide) 20 subunits of insect steroid receptors, or fragments thereof, or insect steroid receptors or fragments thereof, or USP polypeptides of insect juvenile hormone receptors or fragments thereof, which compounds are capable of binding to said receptors which have the effects of either inactivating the receptors (and thus acting as antagonists) or potentiating the activity of the receptor.

25

By "derived from" it is meant that the compounds are based on the three dimensional structure of the aforementioned proteins, that is, synthesized to bind, associate or interfere with insect steroid binding or juvenile hormone binding.

30 The compounds may bind strongly or irreversibly to the ligand binding site or another region of the receptor or USP and act as agonists or antagonists of insect steroids, or juvenile

- 46 -

hormone binding, or otherwise interfere with the binding of ligand, such that ecdysteroids or juvenile hormones. Such compounds would have potent insecticidal activity given the key role of insect steroids, or juvenile hormone, in insect physiology and biochemistry. Such compounds would also possess a unique specificity.

5

This invention is also described with reference to the following non-limiting examples.

EXAMPLE 1

10 Construction of a plasmid (pSV40-EcR) expressing the EcR polypeptide subunit of the D. melanogaster ecdysone receptor

A 3110 base-pair Fsp1-HindIII fragment was excised from a cDNA encoding the EcR polypeptide subunit of the D. melanogaster ecdysone receptor (Koelle et al., 1991), the excised 15 sequence comprising the complete 2634 base pair coding region and 214 base pairs of 5'leader sequence and 258 base pairs of 3'- untranslated sequence. The fragment was ligated into the BamH1 site of the expression plasmid pSG5 (Greene et al, 1988) to produce the expression plasmid pSV40-EcR, wherein expression of the EcR polypeptide subunit of the Drosophila melanogaster ecdysone receptor is placed operably under the control of the SV40 20 promoter sequence.

EXAMPLE 2

Construction of the reporter plasmid p(EcRE),-CAT

25 The reporter plasmid p(EcRE)-CAT was constructed by insertion of seven copies of the hsp27 ecdysone response element, containing a central 13 base pair palindromic ecdysone response element (EcRE), derived from the hsp27 gene (Riddihough and Pelham, 1987) into the HindIII site of the plasmid pMMTV-CAT (Hollenberg and Evans, 1988), 93 base pairs upstream of the transcription start site of the MMTV promoter, thereby operably connecting expression of the 30 chloramphenicol acetyltransferase structural gene to regulation by an insect receptor which binds to the hsp27 ecdysone response element.

- 47 -

EXAMPLE 3

Cell Culture and Transient Transfection

Chinese hamster ovary (CHO) cells were maintained in 50% (v/v) Dubbecco's modified Eagle's medium (DMEM) and 50% (v/v) Hamm F12 nutrient mixture (GIBCO) supplemented with 10% (v/v) foetal bovine serum. Transfection was carried out by the DNA-calcium phosphate coprecipitation method (Ausubel *et al.*, 1992). One day before transfection with the plasmids described in Examples 1 and/or 2, or other expression plasmids, CHO cells were plated out at 5 - 8 x 10⁵ cells per 6 cm diameter culture dish in the above DMEM/F12 medium. Three hours before the addition of the DNA-calcium phosphate co-precipitate, the cells were washed with phosphate buffered saline (PBS; Sambrook *et al.*, 1989) and cultured in fresh DMEM plus 10% (v/v) foetal bovine serum. The cells were incubated in the presence of the co-precipitate for eighteen hours before excess DNA was removed by washing with PBS. The cells were then cultured for another day in DMEM/F12 supplemented with 10% (v/v) foetal bovine serum with or without added ponasterone A (PNA), before harvesting. Cells were washed with PBS, harvested by mechanical scraping in 0.25 M Tris-HCI (pH 7.8), and disrupted by three freeze-thaw cycles.

All transfections included, in addition to expression and reporter plasmids, a β-galactosidase-20 expressing plasmid designated pPgK-LacZ (McBurney *et al*, 1991), which served as an internal control for the efficiency of transfection, and pUC18 DNA in an amount sufficient to produce 10 μg total DNA per culture dish.

The chloramphenical acetyltransferase (CAT) and β-galactosidase activities encoded by the reporter genes present in the reporter plasmids were assayed as described in Sambrook et al, (1989). Cells that were co-transfected with p(EcRE)_T-CAT and pSV40-EcR clearly showed induction of CAT activity in the presence of PNA, showing 50 units of activity. Controls showed negligible activity.

30 We have observed that the ecdysone receptor can lead to stimulation of expression from an ecdysone responsive promoter in some cell types, for example in CHO cells, but not in CV-1

P:\OPER\MRO\ECDYSONE.CIP - 1/7/99

- 48 -

cells. Whilst not being bound by any theory or mode of action, this may reflect a cell-type specific distribution of at least one other transcription factor essential for ecdysone responsiveness. To determine cell types suitable for expressing reporter genes under the control of the steroid receptor of the present invention, the cell-type specificity of ecdysone5 responsive gene expression is assayed in cell-free transcription lysates derived from several target cell lines. Additionally, by fractionating and/or isolating the nuclear proteins of cell lines that express the reporter genes and supplementing lysates derived from non-expressing cell lines with such nuclear protein fractions or isolated proteins, any essential auxiliary factors are defined and the genes encoding them cloned. Co-transfection of the receptor-encoding genes with genes encoding such auxiliary factors removes limitations imposed by cell-type restricted ecdysone responsiveness.

EXAMPLE 4

Testing the Effect of temperature on transient expression

To determine whether the *D. melanogaster* ecdysone receptor polypeptide is stable at physiological temperatures above about 30°C, CHO cells were transfected as described in Example 3, with the plasmid pSV40-EcR and the reporter plasmid p(EcRE)₇-CAT in the 20 presence of PNA, at 30°C and 37°C.

Briefly, CHO cells were plated out at 37°C sixteen to twenty hours before transfection. After washing away the DNA, the cells were cultured for two hours in fresh medium with or without hormone and the dishes divided into duplicate sets. One set was cultured for another day at 37°C before harvesting for CAT and β-galactosidase assays. The other set was cultured for three days at 30°C before assaying enzyme activities. Results indicated a reduction in the fold-induction of gene expression regulated by the *D. melanogaster* ecdysone receptor polypeptide at 37°C, compared to the fold-induction at 30°C, as shown in Table 1.

- 49 -

EXAMPLE 5

Attempted screening of an *L. cuprina* genomic DNA library to isolate genes encoding the EcR polypeptide subunit of the *L. cuprina* ecdysone receptor

- 5 A 627 bp Eco Kpn I fragment encompassing the DNA-binding domain of the EcR polypeptide subunit of the *D. melanogaster* ecdysone receptor was isolated, radioactively labelled and used to screen a *L. cuprina* genomic library constructed in bacteriophage lambda (prepared by CSIRO, division of Entomology, Canberra, Australia). In the first round of screening, twentyfour regions of the plates showed potential positive hybridization to the *D. melanogaster* probe.
- 10 However, second-round screening of these 24 first round positive plaques failed to yield any plaque giving a reproducible positive signal when hybridized to the *D. melanogaster* probe.

TABLE 1

15

pSV40-EcR (µg/dish)	РNA (µМ)	Fold-induction of expression	
		37°C	30°C
2.5	20	14X	35X
	100	59X	54X
0.5	20	8X	26X
	100	47X	33X
0.1	20	1,6X	25X
	100	9.0X	39X

20

EXAMPLE 6 Cloning and characterization of a cDNA molecule encoding the EcR polypeptide of the *L. cuprina* ecdysone receptor

25 Rationale for amplification primer design

The nucleotide sequences of the primers Rdna3 (400>15) and Rdna4 (<400>16) were derived from the amino acid sequence conserved between the DNA-binding domains of the EcR polypeptide subunits of the *D. melanogaster* and *C. tentans* ecdysone receptors. However,

- 50 -

amino acid sequences homologous to two other members of the steroid receptor superfamily of D. melanogaster, Drosophila hormone receptor 3 (DHR3; Koelle, et al., 1991) and Drosophila early gene (E75; Segraves and Hogness, 1990) were excluded from the primer designs, to reduce the possibility of amplifying the L. cuprina homologues of genes encoding 5 DHR3 and/or E75 by PCR.

Amplification primers and PCR conditions

A 105 base pair DNA fragment, encoding the DNA-binding domain of the EcR polypeptide subunit of the L. cuprina ecdysone receptor, was amplified from the L. cuprina genome by 10 PCR, by using the following degenerate primers:

Rdna3 (32mer with EcoRI site):

5'-CGGAATTCCGCCTCTGGTTA(C/T)CA(C/T)TA(C/T)AA(C/T)GC 3' (i.e. <400>15); and Rdna4 (32mer with BamHI site):

15 5'-CGCGGATCC(G/A)CACTCCTGACACTTTCG(C/T)CTCA 3' (i.e. <400>16).

Amplification reactions employed Taql DNA polymerase (Promega) and the following amplification conditions:

cycle 1: 97°C/5 minutes, 50°C hold; add polymerase 50°C/5 minutes;

20 cycles 2-3: 72°C/3 minutes, 94°C/1 minute, 50°C/1 minute;

cycles 4-43: 72°C/3 minutes, 94°C//1 minute, 55°C/1 minute;

cycle 44: 72°C/10 minutes.

To facilitate cloning of the amplified fragments for use as hybridisation probes, the 5' end of 25 primer Rdna3 contained an EcoRI site and the 5' end of primer Rdna4 contained a BamHI site. The amplified L. cuprina gene fragments were cloned into pBluescript SK+, following digestion using the enzymes EcoRI and BamHI, purification of the digested DNA by agarose gel electrophoresis and electro elution of the product band.

30 Hybridisation probe preparation

For probe preparation, the insert was cut out of the pBluescript SK+ vector using EcoR1 and

- 51 -

BamHI, and ³²P-labelled using the GIGAprime DNA Labelling Kit (Bresatec Limited, Adelaide, Australia) essentially according to the manufacturer's instructions, except that random primers were replaced with the specific primers Rdna3 and Rdna4 (see above). Unincorporated label was removed by size exclusion chromatography over Biogel-P60 (Biorad Ltd, Sydney, 5 Australia). The probe was used at 10⁶ cpm/ml in hybridizations.

Construction and screening of L. cuprina cDNA libraries

Two independent *L. cuprina* cDNA libraries derived from late third instar *L. cuprina* larvae were prepared by random priming and oligo-dT priming respectively, and cloned into the *EcoRI* site of the *Lambda/ZapII* vector (Stratagene). The primary libraries generated were subsequently amplified according to the manufacturer's instructions, using standard protocols.

Both cDNA libraries generated are superior to existing *L. cuprina* libraries in terms of their phage titre (i.e. pfu/ml) and insert sizes (0.5 - 4 kbp in both cases). In particular, the primary oligo-dT primed library comprised 4.7 x 10⁶ pfu, whilst the amplified oligo-dT primed library comprised 7.5 x 10¹⁰ pfu/ml; the primary random-primed library comprised 1.3 x 10⁶ pfu, whilst the amplified random-primed library comprised 3.4 x 10¹⁰ pfu/ml.

The prepared cDNA libraries were screened by lifting 500,000 plaques from each library in duplicate on to Hybond N membranes (Amersham) and hybridizing same under low stringency conditions to the ³²P-labelled amplification product produced using the primers Rdna3 and Rdna4 (see above). In particular, hybridisations were performed for twenty four hours at 37°C in a hybridisation solution comprising 42% (w/v) formamide; 5 x SSPE solution; 5 x Denhardt's solution; and 0.1% (w/v) sodium dodecyl sulphate, as described essentially by Ausubel *et al*, (1992) and/or Sambrook *et al*. (1989). The membranes were then washed at 37°C in 2XSSC solution containing 0.1% (w/v) sodium dodecyl sulphate. Following washing, positive plaques were detected by autoradiography, using XOMAT-AR film (Kodak) for two to three days, at -70°C.

30 Two positive-hybridising plaques were obtained from screening of the random-primed library (containing cDNA inserts comprising 561 base pairs and 1600 base pairs in length,

- 52 -

respectively), and one positive-hybridising plaque was obtained from the screening of the oligo-dT primed library (containing a cDNA insert comprising approximately 3400 base pairs in length). pBluescript phagemids containing cDNA inserts were excised *in vivo* from these positive plaques using the Exassist Helper Phage system (Stratagene).

5

The nucleotide sequences of the isolated cDNA clones were obtained using the USB Sequenase Version 2.5 Kit. Sequence data obtained indicated that the 561 bp and 1600 bp cDNAs encode amino acid sequences comprising the important DNA-binding domain and the hormone-binding domain of the EcR polypeptide subunit of the *L. cuprina* ecdysone receptor, whilst the 3400 bp cDNA comprises an entire 2274 bp open reading frame encoding the EcR polypeptide subunit of the *L. cuprina* ecdysone receptor. Accordingly, the 3400 bp cDNA is a full-length cDNA clone. The nucleotide sequence of the open reading frame and 3′-untranslated region is set forth herein as <400> 1. The derived amino acid sequence of the EcR polypeptide subunit of the *L. cuprina* ecdysone receptor encoded by this open reading frame is set out in <400> 2.

EXAMPLE 7

First attempt at cloning and characterization of a cDNA molecule encoding the EcR polypeptide of the *M. persicae* ecdysone receptor

20

Direct screening of a *M. persicae* cDNA library was not effective in isolating a full-length cDNA encoding the EcR polypeptide of the *M. persicae* ecdysone receptor.

DNA encoding the DNA-binding domain of the EcR polypeptide of the *M. persicae* ecdysone receptor was isolated successfully, by amplification as described in Example 6 for the amplification of the homologous *L. cuprina* fragment. The amplified DNA was cloned into pBluescript SK+ and the nucleotide sequence of the cloned insert was obtained using the USB Sequenase version 2.0 Kit, as described in Example 6.

30

Based upon the nucleotide sequence of the amplified DNA fragment, two authentic primers

20

P:\OPER\MRO\ECDYSONE,CIP - 1/7/99

- 53 -

were synthesized as follows:

5'- GCCTCGGGGTATCACTATAACGC -3' (i.e. <400>17); and Mdna1 (23mer):

5'- GCACTCCTGACACTTTCGTCTCA -3' (i.e. <400>18). Mdna2 (23mer):

Hybridisation probe preparation

For M. persicae probe preparation, the amplified 105 bp DNA insert was excised from the pBluescript SK+ vector using EcoRI and BamHI, and 32P-labelled using the GIGAprime DNA Labelling Kit (BresaGen Limited, Adelaide, Australia) essentially according to the 10 manufacturer's instructions, except that random primers were replaced with the specific primers Mdna1 and Mdna2 (see above). Unincorporated label was removed by size exclusion chromatography over Biogel-P60 (Biorad Ltd, Sydney, Australia). The probe was used at 10⁶ cpm/ml in hybridizations.

15 Construction and screening of M. persicae cDNA libraries:

Two independent M. persicae cDNA libraries derived from late third instar M. persicae larvae were prepared by random priming and oligo-dT priming respectively, and cloned into the EcoRI site of the Lambda/Zapl/ vector (Stratagene). The primary libraries generated were subsequently amplified according to the manufacturer's instructions, using standard protocols.

Both cDNA libraries generated are superior to existing M. persicae libraries in terms of their phage titre (i.e. pfu/ml) and insert sizes (0.5 - 4 kbp in both cases). In particular, the primary oligo-dT-primed library comprised 1 x 107 pfu, whilst the amplified oligo-dT primed library comprised 1 x 10¹⁰ pfu/ml; the primary random-primed library comprised 1 x 10⁶ pfu, whilst the 25 amplified random-primed library comprised 2 x 10¹¹ pfu/ml.

Additionally, a further cDNA library was produced in the Lambda ZAP Express insertion vector (Stratagene). To produce this library, cDNA derived from late third instar M. persicae larvae was prepared by oligo-dT priming and cloned directionally into EcoRI-Xhol digested vector 30 DNA. The primary library comprised 1 x 106 pfu, whilst the amplified oligo-dT primed library comprised 1 x 109 pfu/ml, with insert sizes in the range 0.5 - >4 kbp.

PNOPER\MRQ\ECDYSONE.CIP + 171/99

- 54 -

The random-primed *M. persicae* cDNA phage library was screened as described in Example 6, using the *M. persicae* hybridisation probe prepared as described above.

A single positive-hybridising plaque was isolated and sequenced according to standard procedures. The nucleotide sequence of this clone is set forth herein as <400>5. This cDNA clone comprises a 585bp protein-encoding sequence which encodes the DNA-binding domain of a EcR polypeptide of a putative *M. persicae* ecdysone receptor. The amino acid sequence encoded by this partial cDNA clone is set forth herein as <400> 6.

10

EXAMPLE 8

Second attempt at cloning and characterization of a cDNA molecule encoding the EcR polypeptide of the *M. persicae* ecdysone receptor

15 Hybridisation probe preparation

Further hybridisation probes specific for the EcR polypeptide of the *M. persicae* ecdysone receptor were generated using PCR from the Lambda ZAPII oligo dT-primed library using primers AP1 and AP2. The forward primer AP1 was designed to anneal to nucleotide sequences of the partial cDNA (<400>5) encoding part of the first zinc finger motif present in the DNA-binding domain. The reverse primer, AP2, was adapted from degenerate primers designed to anneal to nucleotide sequences complementary to those encoding an EcR ligand binding domain (Kamimura *et al.*, 1996). The nucleotide sequences of primers AP1 and AP2 are as follows:

25 Primer AP1: 5'- TCGTCCGGTTACCATTACAACGC -3' (<400>19); and

Primer AP2; 5'-TAGACCTTTGGC(A/G)AA(C/T)TC(A/G/C/T)ACAAT -3'(<400>20)

The PCR reaction mixture contained 4 µl of each primer (50 pm/µl), 5 µl of deoxynucleotide triphosphate mix (2mM), 1 µl of aphid oligo dT primed Lambda ZAPII cDNA library, 1 µl of 30 recombinant *Pfu* DNA Polymerase (5 units/µl, Stratagene®), 5 µl of 10x *Pfu* buffer (Stratagene®) and 30 µl of MilliQ water. The *Pfu* polymerase was used in this reaction

because it possesses proof-reading activity, which reduces the possibility of misincorporation of nucleotides. The PCR conditions included 42 cycles, each cycle comprising annealing at 55°C, extension at 72°C and melting at 94°C.

5 The major amplification product obtained in this reaction was gel-purified, kinased and ligated into the Smal site of pUC18.

To screen M. persicae cDNA libraries, the cloned amplification product was digested to generate two non-overlapping probes. designated "EcR probe 1" (i.e. <400>7) and "EcR 10 probe 2" (i.e. <400>8). In this regard, digestion of the cloned product with SphI produced a DNA fragment comprising a nucleotide sequence specific for a region encoding the DNAbinding domain (EcR probe 1; <400> 7), whilst digestion with Sphl/EcoRl produced a DNA fragment comprising a nucleotide sequence having homology to a region encoding a putative linker domain, designated domain D, and the 5'-end of a putative hormone-binding domain, 15 present in the EcR polypeptide of the insect ecdysone receptors (EcR probe 2, <400> 8).

EcR probe 1 and EcR probe 2 were labelled with $[\alpha^{-32}P]dATP$ in a reaction catalysed by Klenow fragment. All reagents were components of a GIGAprime DNA labelling kit (BresaGen Limited, Adelaide, Australia), except that the random primers were replaced with specific 20 oligonucleotides synthetisezed to be complementary to the ends of EcR probe 1 and EcR probe 2.

Screening of M. persicae cDNA libraries

480,000 plaques from the oligo dT primed Lambda Zap Express cDNA library (Example 7) 25 were screened as described above, using EcR probe 1. This approach yielded about 300 positive clones. Positive-hybridising clones were pooled and rescreened separately using EcR probe 1 and EcR probe 2, on duplicate lifts. Only four plaques were identified which hybridised to both probes. One of these was found by sequencing to contain a full-length cDNA encoding the EcR polypeptide of the M. persicae ecdysone receptor. The nucleotide sequence of the 30 open reading frame of this cDNA is set forth herein as <400> 9. The derived amino acid sequence of the EcR polypeptide subunit of the M. persicae ecdysone receptor encoded by

P:\OPER\MRO\ECDYSONE.CIP - 1/7/99

- 56 -

this open reading frame is set out in <400> 10.

EXAMPLE 9

In vivo function of recombinant EcR polypeptides of the L. cuprina ecdysone receptor

Construction of plasmid pF3

Plasmid pF3 was constructed in four steps as follows:

First, plasmid p5S1, comprising the full-length cDNA encoding the EcR polypeptide of the L. 10 cuprina ecdysone receptor , was digested with Earl and a 3' Earl cDNA fragment thus generated, encoding the C-terminal end of the EcR polypeptide of the L. cuprina ecdysone receptor, was end-filled and sub-cloned into the HindlI site of pUC19, to construct plasmid pEAR. In plasmid pEAR, the 3' end of the cDNA was oriented towards the KpnI site of the pUC19 vector.

15

Second, plasmid p5S1 was also digested separately with:

- (1) Apol and Pst1, to isolate the 5' end of the cDNA as a 179 bp fragment (fragment A);
- (2) Pstl and Spel, to isolate a 1650 bp cDNA fragment (fragment B); and
- (3) Spel and Bg/II, to isolate a 203 bp fragment (fragment C).

20

Third, plasmid pEAR was digested with Bg/II and KpnI, to isolate the 3' end of the cloned cDNA fragment therein as a 313 bp fragment (fragment D).

Fourth, DNA fragments A, B, C and D were each isolated by agarose electrophoresis and 25 ligated together into pBluescriptSK+, which had been digested with EcoRI and KpnI, to produce plasmid pF3.

Plasmid pF3 thus contains the complete open reading frame of the cDNA encoding the EcR polypeptide of the L. cuprina ecdysone receptor, as a 2368 bp fragment located between two 30 BamHI sites.

P:\OPER\MRO\ECDYSONE.CIP - \(\frac{17}{99}\)

- 57 -

Construction of plasmid pSGLcEcR and plasmid pLcK8

Plasmid pSGLcEcR was constructed by cloning the 2368 bp BamHI fragment from pF3, into the BamHI site of the mammalian expression vector pSG5 (Stratagene). Plasmid pLcK8 is a clone of pSGLcEcR.

5

Construction of plasmid pSGDmEcR

Plasmid pSGDmEcR is identical to plasmid pSV40-EcR (Example 1) comprising the EcR polypeptide of the D. melanogaster ecdysone receptor placed operably under control of the SV40 promoter.

10

15

20

Transfection of CHO cells

CHO cells were co-transfected with a mixture comprising the following DNAs, lysed and assayed for CAT and β-galactosidase enzyme activity, as described in the preceding Examples:

- (1) one of the expression plasmids designated pSGDmEcR, or pSGLcEcR, or the parental expression plasmid pSG5 as a negative control, at a concentration of 1 µg/ml; and
 - (2) the CAT reporter plasmid p(EcRE), CAT at a concentration of 1 µg/ml; and
- (3) an independent LacZ reporter plasmid, pPGKLacZ, at a concentration of 1ug/ml, included as a control to monitor transfection efficiency.

CAT reporter gene expression was induced with 10 µM or 50 µM Muristerone A. In control samples, cells received only the carrier ethanol in place of Muristerone A.

ELISA was used to quantify the synthesis of CAT and β-galactosidase enzymes, in extracts 25 of cells forty eight hours after transfection. Account was taken of the variation between experiments, by normalizing the level of CAT enzyme to the level of β-galactosidase enzyme present in the same extract. Fold induction represents the normalized values for CAT gene expression in cells transfected with pSGDmEcR, pSGLcEcR or pSG5 in the presence of hormone divided by the normalized values for CAT gene expression in cells transfected with 30 the same plasmid but in the absence of hormone. The average values of three independent experiments are shown in Figure 1 and the error bars indicate standard error of the mean.

Data shown in figure 1 indicate that the EcR polypeptide of the *L. cuprina* ecdysone receptor from Example 3 is biologically active *in vivo*. CAT induction is observed at both 50 µM and 10 µm steroid (Muristerone A), with about 30 and 15 fold induction respectively. In view of the *in vivo* activity of the EcR polypeptide of the *L. cuprina* ecdysone receptor obtained according to this protocol, potential insecticidal substances acting by interaction with an insect steroid receptor, such as an ecdysone receptor, are screened by addition of the substances to the *in vivo* assay described herein. Substances are added in an amount from 0.05 µM to 100 µM. Candidate insecticidal compounds are identified by their ability to modulate the reporter gene expression which results from trans-activation by the EcR polypeptide of the *L. cuprina* ecdysone receptor.

EXAMPLE 10

Chimeric EcR polypeptides of insect ecdysone receptors

Chimeric ecdysone receptors comprsing regions derived from EcR polypeptides of ecdysone receptors of different species are produced and assayed for enhanced activity. In a particularly preferred embodiment, a chimeric ecdysone receptor is produced using the EcR polypeptides of the *D. melanogaster*, *M. persicae* and *L. cuprina* ecdysone receptors.

In one exemplification of this embodiment, plasmids pSGLD and pSGDL are produced comprising coding regions derived from the EcR polypeptides of the *D. melanogaster* and *L. cuprina* ecdysone receptors. In plasmid pSGLD, the 5'-end of the open reading frame of the *D. melanogaster* sequence, encoding the N-terminal portion of the EcR polypeptide of the *D. melanogaster* ecdysone receptor to the end of the DNA-binding domain of said polypeptide, is fused to the 3'-end of the open reading frame of the *L. cuprina* sequence, encoding the C-terminal portion of the EcR polypeptide of the *L. cuprina* ecdysone receptor, from the D domain and hormone-binding domain to the carboxyl terminus. In plasmid pSGDL, the 5'-end of the open reading frame of the *L. cuprina* sequence, encoding the N-terminal portion of the EcR polypeptide of the *L. cuprina* ecdysone receptor to the end of the DNA-binding domain of said polypeptide, is fused to the 3'-end of the open reading frame of the *D. melanogaster* sequence, encoding the C-terminal portion of the EcR polypeptide of the *D. melanogaster* ecdysone receptor, from the D domain and hormone-binding domain to the carboxyl terminus.

- 59 -

These plasmids thus encode chimeric EcR polypeptides which form ecdysone receptor variants.

As shown in Figure 2, chimeric EcR polypeptides of *L. cuprina* and *D. melanogaster* ecdysone receptors, comprising fusion polypeptides between the DNA-binding domains and hormone-binding domains of the base *L. cuprina* and *D. melanogaster* polypeptides, exhibit bioactivity when measured in the CAT assay described above. Significant bioactivity of the chimeric EcR polypeptides encoded by plasmids pSGLD and pSGDL, comparable to the bioactivity of the *D. melanogaster* base EcR polypeptide, is observed at both 10 μM and 50 μM concentrations of Muristerone A.

EXAMPLE 11

Isolation and characterisation of a full-length cDNA encoding the EcR partner protein (USP polypeptide) of the *L. cuprina* ecdysone receptor

15 The EcR partner protein (USP polypeptide) subunit of the L. cuprina ecdysone receptor also functions alone as a USP polypeptide of the L. cuprina juvenile hormone receptor. A cDNA encoding both receptor polypeptide activities was isolated using PCR and hybridisation as follows.

20 Hybridisation probe preparation

A 150 base-pair probe, specific for genetic sequences encoding the EcR partner protein (USP polypeptide) subunit of insect ecdysone receptors and/or the USP polypeptide subunit of insect juvenile hormone receptors (<400>13), was isolated by PCR from *L. cuprina* genomic DNA using the degenerate primers described by Tzertzinis *et al.* (1994). The PCR reaction conditions were as described in Example 6, except that *Pfu* polymerase was used in place of *Taql* polymerase.

The amplified DNA fragment was sub-cloned into *EcoRI* and *Clal* double-digested *p*Bluescript SK+ vector (Stratagene), after double-digestion of the fragment using the enzymes *EcoRI* and 30 *Clal*, purification of the amplified fragment by agarose gel electrophoresis, and electro elution of the product band. The nucleotide sequence of the probe was obtained using the USB

- 60 -

Sequenase version 2.0 Kit (<400> 13).

For probe preparation, the amplified L. cuprina DNA fragment was excised from the vector using EcoRI and Sall, gel purified and ³²P-labelled using the GIGAprime DNA Labelling Kit 5 (BresaGen Limited, Adelaide, Australia) essentially according to the manufacturer's instructions except that random primers were replaced with the two degenerate primers described by Tzertzinis et al. (1994) (see above). Unincorporated label was removed by size exclusion chromatography over Biogel-P60 (Biorad Ltd, Sydney, Australia). The probe was used at 106 cpm/ml in hybridizations.

10

20

Screening of L. cuprina cDNA libraries

The L. cuprina cDNA library described above (Example 6) was screened with the amplified probe as described in Example 6. The nucleotide sequence of the full-length open reading frame of this cDNA molecule and amino acid sequence therefor, are set forth herein as <400> 15 3 and <400> 4, respectively.

EXAMPLE 12

Isolation and characterisation of a partial cDNA encoding the EcR partner protein (USP polypeptide)of the M. persicae ecdysone receptor

The EcR partner protein (USP polypeptide) subunit of the M. persicae ecdysone receptor also functions alone as a USP polypeptide of the M. persicae juvenile hormone receptor. To isolate a partial cDNA encoding both receptor polypeptide activities, a 140 bp probe was amplified 25 from M. persicae genomic DNA, by PCR, using the two degenerate primers described by Tzertzinis et al.(1994) (see preceding Example). The PCR reaction conditions were as described in Example 6, except that Pfu polymerase was used in place of Taql polymerase.

The amplified DNA fragment was sub-cloned into EcoRI and Clal double-digested pBluescript 30 SK+ vector (Stratagene), after double-digestion of the fragment using the enzymes EcoRI and Clal, purification of the amplified fragment by agarose gel electrophoresis, and electro elution

- 61 -

of the product band.

The nucleotide sequence of the insert in the pBluescript SK+ vector was obtained using automated fluorescent dye terminator sequencing (SUPAMAC, Sydney Australia).

5

Hybridisation probe preparation and library screening

For probe preparation the amplified M. persicae DNA insert was cut out of the pBluescript+ vector with EcoRI and Sall, gel purified and 32P-labelled using the GIGAprime DNA Labelling Kit (Bresatec Limited, Adelaide, Australia) essentially according to the manufacturer's 10 instructions except that random primers were replaced with the degenerate primers described by Tzertzinis et al.(1994) (see preceding Example). Unincorporated label was removed by size exclusion chromatography over Biogel-P60 (Biorad Ltd, Sydney, Australia). The probe was used at 10⁶ cpm/ml in hybridizations to screen the M. persicae cDNA library as described in Examples 7 and 8.

15

The positive-hybridising clones were plaque-purified and sequenced using standard procedures as described herein. The nucleotide sequence of the open reading frame of the full-length cDNA encoding the partner protein (USP polypeptide) subunit of the M. persicae ecdysone receptor or the USP polypeptide of the M. persicae juvenile hormone receptor is set 20 forth herein as <400> 11. The derived amino acid sequence of this open reading frame is set forth as <400>12.

EXAMPLE 13

A construct for the baculovirus-directed co-expression of functional ligand-binding regions of the EcR polypeptide and partner protein (USP polypeptide) of the D. 25 melanogaster ecdysone receptor

A vector was prepared to facilitate the baculovirus-directed co- expression of ligand-binding regions derived from the EcR polypeptide and partner protein (USP polypeptide) of the D. 30 melanogaster ecdysone receptor, the protein products of which associate on co-expression to form a functional hormone-binding complex. The associated proteins are then used in high

- 62 -

through-put assays or three-dimensional structural analysis. We have found that the ligandbinding domain, together with most of the linker domain of the EcR polypeptide subunit and of the EcR partner protein (USP polypeptide), are ssufficient to associate to form a functional hormone-binding complex.

5

1. Isolation of the ligand-binding region and linker region of the EcR polypeptide of the D. melanogaster ecdysone receptor.

A Sac I- HindIII fragment encoding most of the linker (domain D) and all of the ligand-binding domain (domains E and F) of the EcR polypeptide of the Drosophila melanogaster ecdysone 10 receptor was excised from a plasmid comprising DNA encoding the complete EcR polypeptide (Koelle et al. 1991). The excised fragment was cloned into Sacl - HindllI-digested expression vector pQE31(Qiagen), to produce the plasmid vector pQE31DmECR.

- 2. Construction of a baculovirus expressing the linker regions of EcR and USP polypeptides 15 A baculovirus was constructed for the co-expression in insect cells of:
 - a cDNA region comprising a nucleotide sequence which encodes at least the (i) ligand-binding domain and much of the linker domain of the EcR polypeptide of the D. melanogaster ecdysone receptor isolated as described at paragraph (1) above; and
- a cDNA region comprising a nucleotide sequence which encodes at least the (ii) 20 ligand-binding domain and much of the linker domain of the partner protein (USP polypeptide) of the D. melanogaster ecdysone receptor.

To produce this baculovirus, a EcoR I - HindlII fragment was excised from pQE31DmECR, said fragment encoding an oligo-His tag, and most of the linker domain, together with all of the 25 ligand-binding domain of EcR polypeptide. This EcoR I - HindIII fragment was ligated into EcoR I - HindIII cleaved pFastBacDUAL, to produce the plasmid pDmEcR.DUAL. To insert gene sequences specific for the partner protein (USP polypeptide), the Hindli - Nsil fragment encoding most of the linker and all of the ligand-binding domain of the partner protein (USP polypeptide) was excised from a full-length cDNA clone in plasmid pZ7-1 (supplied by Vince 30 Henrich) and ligated into Ncol - Nsil cleaved pDmEcR.DUAL. A nucleotide sequence encoding a "FLAG" peptide was subsequently incorporated upstream of, and in the same reading frame

- 63 -

as, the nucleotide sequence encoding the linker and ligand-binding regions of the partner protein (USP polypeptide), by ligation into the unique Smal site, thereby producing the plasmid pDmEcR.USP.DUAL. Plasmids containing the FLAG-encoding nucleotide sequence in the correct orientation were selected by nucleotide sequence determination.

5

The segment of pDmEcR.USP.DUAL which encodes the tagged linker and ligand-binding regions of the EcR polypeptide and partner protein (USP polypeptide) sequences, placed operably under the control of polyhedrin and p10 promoters, respectively, was recombined into a baculovirus genome, by employing the Tn7 transposition system (Luckow et al, (1993). The 10 polypeptide products were then co-expressed in insect Sf21 and Sf9 cells, where they associated into a functional complex.

Expression of the tagged linker and ligand-binding regions of the EcR polypeptide and partner protein (USP polypeptide) sequences was examined by immunoblot analysis of extracts 15 derived from insect Sf21 cells infected with the recombinant baculovirus, employing antibodies directed against the oligo-His and FLAG tags. This analysis detected bands on immunoblot analysis of approximately the predicted sizes for the expressed tagged linker and ligandbinding regions of the EcR polypeptide and partner protein (USP polypeptide).

20 The protein detected by anti-oligo-His-antibodies was enriched by affinity purification on nickel-NTA resin (Qiagen), and the FLAG-labelled protein was affinity-purified using FLAG M2 Affinity Gel (Kodak). It was further demonstrated that the oligo-His-tagged EcR polypeptide and the FLAG-tagged EcR partner protein (USP polypeptide) bound as a hetero-oligomeric complex to FLAG M2 Affinity Gel (Kodak).

25

Furthermore, binding assays, performed using a modification of the method of Yund et al (1978), demonstrated a highly-significant increase in the binding of the a labelled ecdysone analogue, [3H]ponasterone A, in cells infected by the recombinant baculovirus, compared to the binding observed for the naturally-occurring ecdysone holoreceptor in L. cuprina embryos. 30 In contrast, cells infected by a control virus displayed neither antibody-positive bands on

western analysis, nor specific binding of [3H] ponasterone A, above background levels. These

- 64 -

data indicate correct folding and association of the variant polypeptides comprising the linker and ligand-binding regions of the *D. melanogaster* EcR polypeptide and *D. melanogaster* partner protein (USP polypeptide). The correctly-folded and associated complex formed by the truncated Ecr polypeptide and trucated EcR partner protein (USP polypeptide), is used for 5 X-ray and NMR structural analysis and for high-throughput screens.

EXAMPLE 14

Construct for the baculovirus-directed co-expression of functional ligand-binding regions of the EcR polypeptide and partner protein (USP polypeptide) of the L.

10 cuprina ecdysone receptor

A vector for the baculovirus-directed co- expression of ligand-binding domains derived from the EcR polypeptide and partner protein (USP polypeptide) of the *L. cuprina* ecdysone receptor was prepared essentially as described in the preceding Example.

15

1. Isolation of the ligand-binding region and linker region of the EcR polypeptide of the L. cuprina ecdysone receptor.

A SphI – KpnI fragment encoding most of the linker (domain D) and all of the ligand-binding domain (domains E and F) of the EcR polypeptide of the L. cuprina ecdysone receptor was excised from a cDNA clone encoding the complete EcR polypeptide and cloned into the SphI – KpnI cleaved expression vector pQE32 (Qiagen), to produce the plasmid pQE32LcEcR.

- 2. Isolation of the ligand-binding region and linker region of the partner protein (USP polypeptide) of the *L. cuprina* ecdysone receptor.
- 25 A DNA fragment encoding most of the linker domain and all of the ligand-binding domain of the partner protein (USP polypeptide) of the L. cuprina ecdysone receptor was sub-cloned to produce the plasmid pBLU1.
- 3. Construction of a baculovirus expressing the linker regions of *L. cuprina* EcR and USP 30 polypeptides

A baculovirus was constructed for the co-expression in insect cells of:

P:\OPER\MRO\ECDYSONE,CIP - 1/7/99

- a cDNA region comprising a nucleotide sequence which encodes at least the (i) ligand-binding domain and much of the linker domain of the EcR polypeptide of the L. cuprina ecdysone receptor isolated as described at paragraph (1) above; and
- (ii) a cDNA region comprising a nucleotide sequence which encodes at least the ligand-binding domain and much of the linker domain of the partner protein (USP polypeptide) of the L. cuprina ecdysone receptor isolated as described at paragraph (2) above.

To produce this baculovirus, a EcoR I - PstI fragment derived from plasmid pQE32LcEcR, 10 encoding an oligo-His tag and most of the linker domain together with all of the ligand-binding domain of the L. cuprina EcR polypeptide was ligated into EcoRI- Pstl cleaved pFastBac.DUAL, to produce the plasmid pLcEcR.DUAL. An Avall-EcoRV fragment, encoding most of the linker and all of the ligand-binding domain of L. cuprina partner protein (USP polypeptide) was excised from plasmid pBLU1 and ligated, together with a "FLAG" encoding 15 sequence into the Pvull site of pLcEcR.DUAL, to produce plasmid pLcEcR.USP.DUAL.

The segment of pLcEcR.USP.DUAL which encodes the tagged linker and ligand-binding regions of the EcR polypeptide and partner protein (USP polypeptide) sequences, placed operably under the control of polyhedrin and p10 promoters, respectively, was recombined into 20 a baculovirus genome, by employing the Tn7 transposition system (Luckow et al., (1993). The polypeptide products were then co-expressed in insect Sf21 and Sf9 cells, where they associated into a functional complex.

Expression was examined by immunoblot analysis. Antibodies directed against oligo-His and 25 FLAG tags detected bands on immunoblot analysis of approximately the predicted sizes for the expressed EcR and USP polypeptide regions respectively, in extracts from insect Sf21 cells infected with the recombinant baculovirus. The protein detected by anti-oligo-His was greatly enriched utilising a nickel-NTA resin (Qiagen) and the FLAG-labelled protein purified on FLAG M2 Affinity Gel (Kodak). It was also demonstrated by immunoblot analysis that oligo-30 His-tagged L. cuprina truncated EcR polypeptides and FLAG-tagged L. cuprina truncated EcR partner protein (USP polypeptide) bind as a hetero-oligomeric complex to FLAG M2 Affinity PAGPERIMROLECDYSONE.CIP - 1/7/99

- 66 -

Gel (Kodak).

Furthermore, binding assays, carried out by a modification of the method of Yund et al. (1978), demonstrated a highly-significant increase in the binding of the tritiated ecdysone analogue, ponasterone A, in cells infected by recombinant virus indicating correct folding and association of the two protein subunits (Figure 3), greater than that of the ecdysone holoreceptor in L. cuprina embryos. Cells infected by a control virus displayed neither antibody-positive bands on western analysis nor specific binding of tritiated hormone above background.

10 Expression of the tagged linker and ligand-binding regions of the *L. cuprina* EcR polypeptide and partner protein (USP polypeptide) sequences was examined by immunoblot analysis of extracts derived from insect Sf21 cells infected with the recombinant baculovirus, employing antibodies directed against the oligo-His and FLAG tags. This analysis detected bands on immunoblot analysis of approximately the predicted sizes for the expressed tagged linker and ligand-binding regions of the *L. cuprina* EcR polypeptide and partner protein (USP polypeptide).

The protein detected by anti-oligo-His-antibodies was enriched by affinity purification on nickel-NTA resin (Qiagen), and the FLAG-labelled protein was affinity-purified using FLAG M2 Affinity 20 Gel (Kodak).

Furthermore, binding assays, performed using a modification of the method of Yund *et al* (1978), demonstrated a significant increase in the binding of the labelled ecdysone analogue, [³H] ponasterone A, in cells infected by the recombinant baculovirus, compared to the binding observed for the naturally-occurring ecdysone holoreceptor in *L. cuprina* embryos (Figure 3). In contrast, cells infected by a control virus displayed neither antibody-positive bands on western analysis, nor specific binding of [³H] ponasterone A, above background levels.

These data indicate correct folding and association of the variant polypeptides comprising the 30 linker and ligand-binding regions of the *L. cuprina* EcR polypeptide and *L. cuprina* partner protein (USP polypeptide). The correctly-folded and associated complex formed by the

P-YOPER\MEO\ECDYSONE CIP . 1/7/99

- 67 -

truncated Ecr polypeptide and trucated EcR partner protein (USP polypeptide), is used for Xray and NMR structural analysis and for high-throughput screens.

EXAMPLE 15

A construct for the expression of the ligand-binding region of the USP polypeptide 5 of the L. cuprina juvenile hormone receptor

The donor plasmid pLcEcR.USP.DUAL (Example 14) was digested with BssHII and PstI to remove the L. cuprina EcR polypeptide-encoding segment therein, thereby leaving the tagged 10 linker and ligand-binding regions of the L. cuprina USP polypeptide-encoding nucleotide sequence. The digested plasmid was blunt-ended using T4 DNA polymerase and Klenow polymerase, isolated by gel purification, and finally re-ligated to produce the plasmid pLc.USP.SINGLE.

15 To produce recombinant baculovirus capable of expressing the tagged linker and ligandbinding regions of the USP polypeptide, the segment of pLc.USP.SINGLE encoding this polypeptide and the p10 promoter sequence to which said segment is operably connected, is recombined into a baculovirus genome employing the Tn7 transposition system (Luckow et al., 1993). The polypeptide product is then expressed to form a functional juvenile hormone-20 binding polypeptide and preferably, a modulator of a juvenile hormone receptor. The correctlyfolded truncated USP polypeptide is used for X-ray and NMR structural analysis and for highthroughput screens.

EXAMPLE 16

In-vitro Screening for the Detection of Insecticidal Compounds

The EcR partner protein (USP polypeptide) of the insect ecdysone receptor and USP polypeptide of the insect juvenile hormone receptor of the present invention, optionally associated with the EcR polypeptides of insect ecdysone receptors of the present invention 30 as described in the preceding Examples, are coupled to pins according to the procedure of Geysen et al. (1987), and reacted with candidate insecticidal compounds, generally at a

- 68 -

concentration in the range from about 0.05 µM to about 100 µM of the candidate compound. The binding of compounds is detected using standard procedures, and compounds having insecticidal activity are identified. Preferably, such compounds exhibit insecticidal activity against a range of insects, including diptera, hemiptera, coleoptera, ants, and moths, amongst others. More preferably, the compounds will exhibit insecticidal activity against *L. cuprina*, *M. persicae*, *D. melanogaster*, scale insect, white fly, and leaf hopper, amongst others. In a particularly preferred embodiment, insecticidal compounds are specific to *L. cuprina* and/or *M. persicae* and close relatives thereof.

10

EXAMPLE 17

Cloned Myzus persicae EcR/USP complex binds ponasterone A in vitro.

In vitro-translated Myzus persicae EcR (MpEcR) polypeptide and an in vitro-translated M.persicae USP (MpUSP) polypeptide were produced labelled with [35S]Methionine, using the Promega TNT-Coupled Reticulocyte Lysate System. Each batch of lysate contained 100-200 mg/ml of endogenous proteins (using BSA as a standard). The products were analysed by SDS-PAGE and radioautography. The results confirmed that the cloned cDNAs encode proteins of the sizes predicted from the length of putative open reading frames of the cDNAs present in plasmids pMpEcR and pMPUSP. The yields of EcR and USP were similar as assessed by SDS-PAGE.

20

In functional assays, DNA plasmids pMpEcR (AGAL Accession No. NM99/04567; 1 mg) or pMpUSP (AGAL Accession No. NM99/04568; 1 mg), which have been constructed using the vector pBK-CMV, and 1 ml of appropriate TNT RNA Polymerase were added to 48 ml of reaction mix which contained TNT Lysate, TNT Reaction Buffer, amino acid mixture, Rnasin Ribonuclease Inhibitor and nuclease-free water in volumes specified in the manufacture ps protocol. In control reactions, a Luciferase T3 control DNA (Promega) was used in place of pMpEcR or pMpUSP. T7 RNA Polymerase was used for transcription of the *M. persicae* EcR RNA from plasmid pMpEcR, whilst T3 RNA Polymerase was used for transcription of *M. persicae* USP RNA from the plasmid pMpUSP and the Luciferase T3 control DNA. The 30 reactions were carried out for 90 minutes at 30°C.

- 69 -

The control reaction produces 150-500 ng of luciferase per 50 ml reaction.

The ecdysteroid binding activities of an *in vitro*-translated *Myzus persicae* EcR (MpEcR) polypeptide and an *in vitro*-translated complex of the *M.persicae* EcR and USP polypeptides were produced from the RNAs using the TNT-Coupled Reticulocyte Lysate System (Promega). The mixtures were stored at -20°C overnight.

After thawing the translation products, 15 ml aliquots of the reaction mixture containing M.persicae EcR and USP polypeptides were combined to promote formation of the EcR/USP 10 complex, For assays of individual proteins, 15 ml of the reaction mixture containing M.persicae EcR polypeptide or 15 ml of the reaction mixture containing M.persicae USP polypeptide was combined with 15 ml of control luciferase protein reaction mixture. Samples were each diluted to 435 ml with EcR40 buffer [40 mM KCl, 25 mM HEPES pH 7.0, 1 mM EDTA, 1mM DTT, BSA(0.5mg/ml), 10% glycerol] to allow for triplicates in the ligand binding assay. A control 15 reaction (Blank) was established which contained EcR40 buffer only. An aliquot (140 ml) of each diluted sample was incubated with tritiated ponasterone A (DuPont NEN, Batch Number 3281108) at a final concentration of 2.2 nM for 90 min at room temperature. After incubation, the ligand binding reactions were placed on ice. The samples were pipetted onto Whatman GF/C filters and incubated for 30 sec. The filters were then placed on a vacuum sinter, washed 20 with 10 ml EcR40 buffer and transferred to scintillation vials. After adding 7 ml of InstaGel Plus to each vial, the contents were vortexed and left at room temperature until the filters became transparent. The receptor bound ligand was quantified using a TriCarb 2100TR scintillation counter.

25 The results depicted in Figure 4 indicate that significantly higher amounts of ponasterone A bind to the complex than to iether the USP or EcR polypeptides alone.

Each reference cited herein is incorporated by reference to the extent that it is not inconsistent with the present disclosure.

- 70 -

REFERENCES

- 1. Amann and Brosius (1985). Gene 40: 183.
- Ausubel, et al (1992), Current Protocols in Molecular Biology, Greene/Wiley, New York 2.
- 3. Bugg, et al (1993) Scientific American, December Issue: pp 60-66.
- 5 4. Devereux, J., et al. (1984). Nucl. Acids Res. 12:387-395.
 - 5, Evans (1988) Science, 240: 889-895.
 - 6. Geysen et al (1987) J. Immunol Methods, 102, 259-274.
 - 7. Greene et al. (1988) Nucleic Acids Res. 15:369.
 - 8. Hollenberd and Evans (1988) Cell 55: 899-906.
- 10 9. Hollenberg et al (1991) Cell, 67, 59-77.
 - 10. Jones, G and Sharp, P, (1997) Proc. Nat. Acad. Sci. USA 94:13499-13503.
 - Karnimura et al (1996) Comp. Biochem. Physiol. 113, 341-347. 11.
 - 12. Koelle et al (1991) Cell, 67:59-77
 - 13. Krust et al (1986) EMBO. J., 5: 891-897.
- 15 14. Luckow et al., (1993) J. Virol, 67: 4566.
 - McBurney et al (1991), Nucleic Acids Res., 19, 5755-5761. 15.
 - 16. McPherson, M.J., Quirke, P. and Taylor, G.R. (1991) PCR A Practical Approach. IRL Press, Oxford University Press, Oxford, United Kingdom.
 - 17. Needleman and Wunsch (1970) J. Mol. Biol. 48:443-453.
- 20 18. Potrykus (1990) Bio/Technology 8: 535-542.
 - 19. Riddihough, G., and Pelham, H. R. B., (1987), EMBO J., 6, 3729-3734
 - 20. Sambrook et al (1989), Molecular Cloning: A Laboratory Manual, Cold Springs Harbor Laboratory Press
 - 21. Segraves and Hogness (1990) Genes Devel. 4: 204-219.
- 25 22. Shimatake and Rosenberg (1981) Nature 292: 128
 - 23. Studier and Moffat (1986) J. Mol. Biol. 189: 113
 - 24. Thompson, et al., (1994) Nucl. Acids Res. 22:4673-4680.
 - 25. Tzertzinis et al (1994) J. Mol. Biol. 258, 479-486.
 - Von Itzstein et al (1993) Nature Vol. 363: 418-423. 26.
- 30 27. Yund, et al. (1978) Proc. Nat. Acad. Sci. USA, 24: 6039-6043.

- 71 -

WE CLAIM:

- An isolated nucleic acid molecule comprising a nucleotide sequence which encodes or is complementary to a sequence which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide or a bioactive derivative or analogue thereof, wherein said polypeptide is selected from the groups consisting of (i) an EcR polypeptide of a steroid receptor; (ii) the partner protein (USP polypeptide) of a steroid receptor; and (iii) the USP polypeptide of a juvenile hormone receptor; and wherein said polypeptide comprises an amino acid sequence that is at least 40% identical to an amino acid sequence selected from the group consisting of:
 - an amino aicd sequence set forth in any one of <400>2, <400>4, <400>6, (a) <400>10, <400>12 or <400>14; and
 - an amino acid sequence encoded by a cDNA present in any one of the (þ) plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568.
- 2. The isolated nucleic acid molecule according to claim 1, wherein the steroid receptor is an ecdysteroid receptor.
- 3. The isolated nucleic acid molecule according to claim 2, wherein the ecdysteroid receptor is an insect ecdysone receptor.
- 4. The isolated nucleic acid molecule according to claim 3, wherein the insect ecdysone receptor comprises the EcR polypeptide of an insect ecdysone receptor or the partner protein (USP polypeptide) of an insect ecdysone receptor.
- 5. The isolated nucleic acid molecule according to claim 4, wherein the insect is selected from the list comprising dipteran, hemipteran, coleopteran, lepidopteran, and neuropteran insects and ants.
- 6. The isolated nucleic acid molecule according to claim 5, wherein the hemipteran insect is Myzus persicae or a close relative thereof.

- The isolated nucleic acid molecule according to claim 6, wherein the insect steroid 7. receptor polypeptide comprises an EcR polypeptide of the M. persicae ecdysone receptor having the amino acid sequence set forth in <400>6 or <400>10 or encoded by the cDNA present in plasmid pMpEcR (AGAL Accession No. NM99/04567) or a bioactive analogue or derivative thereof.
- The isolated nucleic acid molecule according to claim 6, wherein the insect steroid 8. receptor polypeptide comprises an EcR partner protein (USP polypeptide) of the M. persicae ecdysone receptor or a USP polypeptide of the M. persicae juvenile hormone receptor having or including the amino acid sequence set forth in <400>12 or encoded by the cDNA present in plasmid pMpUSP (AGAL Accession No. NM99/04568) or a bioactive analogue or derivative thereof.
- 9. The isolated nucleic acid molecule according to claim 5, wherein the dipteran insect is L. cuprina or a close relative thereof.
- The isolated nucleic acid molecule according to claim 9, wherein the insect steroid 10. receptor polypeptide comprises an EcR polypeptide of the L. cuprina ecdysone receptor having the amino acid sequence set forth in <400>2 or encoded by the cDNA present in plasmid pLcEcR (AGAL Accession No. NM99/04566) or a bioactive analogue or derivative thereof.
- 11. The isolated nucleic acid molecule according to claim 9, wherein the insect steroid receptor polypeptide comprises an EcR partner protein (USP polypeptide) of the L. cuprina ecdysone receptor or a USP polypeptide of the L. cuprina juvenile hormone receptor having the amino acid sequence set forth in <400>4 or <400>14 or encoded by the cDNA present in plasmid pLcUSP (AGAL Accession No. NM99/04565) or a bioactive analogue or derivative thereof.
- 12. The isolated nucleic acid molecule according to claim 1, wherein the bioactive derivative or analogue comprises a fragment of an EcR polypeptide of an insect ecdysone receptor or a fragment of an EcR partner protein (USP polypeptide) of an insect ecdysone

receptor, wherein said fragment includes at least one ligand-binding region of said EcR polypeptide or said EcR partner protein (USP polypeptide).

- 13. The isolated nucleic acid molecule according to claim 12, wherein the ligand-binding region comprises a linker domain of the EcR polypeptide or a linker domain of the EcR partner protein (USP polypeptide).
- 14. The isolated nucleic acid molecule according to claim 12, wherein the ligand-binding region comprises a hormone-binding domain of the EcR polypeptide or a hormone-binding domain of the EcR partner protein (USP polypeptide).
- 15. The isolated nucleic acid molecule according to claim 12, wherein the ligand-binding region comprises a linker domain and hormone-binding domain of the EcR polypeptide or a linker domain and hormone-binding domain of the EcR partner protein (USP polypeptide).
- 16. The isolated nucleic acid molecule according to claim 1, comprising a protein-encoding nucleotide sequence which is at least 40% identical to any one of the nucleotide sequences set forth in <400>1, <400>3, <400>5, <400>9, <400>11 or <400>13 or a complementary nucleotide sequence thereto or the cDNA present in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568.
- 17. An isolated nucleic acid molecule comprising a nucleotide sequence which encodes or is complementary to a sequence which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide or a bioactive derivative or analogue thereof, wherein said nucleotide sequence is selected from the list comprising:
 - (i) a nucleotide sequence having at least 40% identity to any one of the nucleotide sequences set forth in <400>1, <400>3, <400>5, <400>9, <400>11 or <400>13 or a complementary nucleotide sequence thereto;
 - (ii) a nucleotide sequence that is capable of hybridising under at least low stringency conditions to any one of the nucleotide sequences set forth in <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11 or <400>13 or to a

complementary nucleotide sequence thereto;

- a nucleotide sequence having at least 40% identity to a nucleotide sequence of a cDNA present in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568;
- a nucleotide sequence that is capable of hybridising under at least low (iv) stringency conditions to a cDNA present in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568; and
- a nucleotide sequence that is amplifiable by PCR using a nucleic acid primer (v) sequence set forth in any one of <400>15, <400>16, <400>17, <400>18, <400>19 or <400>20.
- 18. An isolated nucleic acid molecule which encodes an insect steroid receptor polypeptide and comprises the nucleotide sequence set forth in <400>1 or a complementary nucleotide sequence thereto or the nucleotide sequence of the cDNA present in plasmid pLcEcR (AGAL Accession No. NM99/04566).
- 19. An isolated nucleic acid molecule which encodes an insect steroid receptor polypeptide or a juvenile hormone receptor polypeptide and comprises the nucleotide sequence set forth in <400>3 or <400>13 or a complementary nucleotide sequence thereto or the nucleotide sequence of the cDNA present in plasmid pLcUSP (AGAL Accession No. NM99/04565).
- 20, An isolated nucleic acid molecule which encodes an insect steroid receptor polypeptide and comprises the nucleotide sequence set forth in <400>5 or <400>7 or <400>8 or <400>9 or a complementary nucleotide sequence thereto or the nucleotide sequence of the cDNA present in plasmid pMpEcR (AGAL Accession No. NM99/04567).
- 21. An isolated nucleic acid molecule which encodes an insect steroid receptor polypeptide or a juvenile hormone receptor polypeptide and comprises the nucleotide sequence set forth in <400>11 or a complementary nucleotide sequence thereto or the nucleotide sequence of the cDNA present in plasmid pMpUSP (AGAL Accession No. NM99/04568).

- A method of identifying an isolated nucleic acid molecule which encodes an insect 22. steroid receptor polypeptide which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide comprising:
 - hybridising genomic DNA, mRNA or cDNA with a hybridisation-effective (i) amount of one or more probes selected from the list comprising:
 - probes comprising at least 10 contiguous nucleotides in length derived (a) from any one of <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11, <400>13, <400>15, <400>16, <400>17, <400>18, <400>19 or <400>20 or a complementary nucleotide sequence thereto;
 - probes comprising at least 10 contiguous nucleotides in length derived (b) from a cDNA contained in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568; and
 - hybridisation probes comprising the nucleotide sequences set forth in any one of <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11, or <400>13 or a complementary nucleotide sequence thereto or a homologue, analogue or derivative thereof which is at least 40% identical to said sequence or complement; and
 - (ii) detecting the hybridisation.
- 23. The method of claim 22 wherein the step of detecting the hybridisation comprises detecting a reporter molecule that is covalently bound to the probe.
- 24. A method of identifying an isolated nucleic acid molecule which encodes an insect steroid receptor polypeptide which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide comprising:
 - annealing to genomic DNA, mRNA or cDNA, one or more PCR primers comprising at least 10 contiguous nucleotides in length derived from the group consisting of:
 - a primer derived from any one of <400>1, <400>3, <400>5, <400>7, (a) <400>8, <400>9, <400>11, <400>13, <400>15, <400>16, <400>17, <400>18, <400>19 or <400>20 or a complementary nucleotide sequence thereto; and

- (b) a primer derived from a cDNA contained in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568; and
- (ii) amplifying a nucleotide sequence which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide in a polymerase chain reaction.
- 25. A method of identifying an isolated nucleic acid molecule which encodes an insect steroid receptor polypeptide which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide comprising:
 - (i) amplifying a nucleotide sequence which encodes a steroid receptor polypeptide or a juvenile hormone receptor polypeptide in a polymerase chain reaction using one or more PCR primers comprising at least 10 contiguous nucleotides in length from the group consisting of:
 - (a) a primer derived from any one of <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11, <400>13, <400>15, <400>16, <400>17, <400>18, <400>19 or <400>20 or a complementary nucleotide sequence thereto; and
 - (b) a primer derived from a cDNA contained in any one of the plasmids deposited under AGAL Accession Nos NM99/04565, NM99/04566, NM99/04567, or NM99/04568;
 - (ii) hybridising the amplified nucleotide sequence to genomic DNA, mRNA or cDNA with a hybridisation-effective amount of one or more probes selected from the group consisting of:
 - (a) a probe comprising at least 10 contiguous nucleotides in length derived from any one of <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11, <400>13, <400>15, <400>16, <400>17, <400>18, <400>19 or <400>20 or a complementary nucleotide sequence thereto;
 - (b) a probe comprising at least 10 contiguous nucleotides in length derived from a cDNA contained in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568; and
 - (c) hybridisation probes comprising the nucleotide sequences set forth in any one of <400>1, <400>3, <400>5, <400>7, <400>8, <400>9, <400>11, or

- <400>13 or a complementary nucleotide sequence thereto or a homologue, analogue or derivative thereof which is at least 40% identical to said sequence or complement; and
- (iii) detecting the hybridisation.
- The method of claim 25 wherein the step of detecting the hybridisation comprises 26. detecting a reporter molecule that is covalently bound to the probe.
- 27. The method according to claim 22, further comprising the step of isolating the identified nucleic acid molecule.
- 28. A genetic construct comprising the isolated nucleic acid molecule according to claim 1 operably linked to a promoter sequence.
- 29. The genetic construct according to claim 28, wherein the promoter is the SV40, MMTV, polyhedron or p10 promoter.
- 30. A recombinant or isolated polypeptide comprising a steroid receptor polypeptide or juvenile hormone receptor polypeptide or a bioactive derivative or analogue thereof, wherein said polypeptide:
 - (i) is selected from the list comprising EcR polypeptide of a steroid receptor, the partner protein (USP polypeptide) of a steroid receptor and the USP polypeptide of a juvenile hormone receptor; and
 - comprises an amino acid sequence that is at least 40% identical to any one of the amino acid sequences selected from the group consisting of:
 - an amino acid sequence set forth in <400>2, <400>4, <400>6, <400>10, <400>12 or <400>14;and
 - an amino acid sequence encoded by a cDNA present in any one of the plasmids deposited under AGAL Accession Nos. NM99/04565, NM99/04566, NM99/04567, or NM99/04568;

wherein said polypeptide is substantially free of naturally-associated cellular components.

- A cell comprising the nucleic acid molecule according to claim 1. 31.
- 32. A cell comprising the genetic construct according to claim 28,
- 33. A cell which expresses the isolated or recombinant polypeptide according to claim 30.
- 34. A method of identifying a modulator of steroid receptor-mediated gene expression or juvenile hormone receptor-mediated gene expression comprising:
 - (ĭ) assaying the expression of a reporter gene in the presence of the recombinant or isolated polypeptide according to claim 30 and a potential modulator; and
 - (ii) assaying the expression of the reporter gene in the presence of the recombinant or isolated polypeptide according to claim 30 and without said potential modulator; and
 - (ii) comparing expression of the reporter gene in the presence of the potential modulator to the expression of a reporter gene in the absence of the potential modulator,

wherein said reporter gene is placed operably under the control of a steroid response element (SRE) to which said steroid receptor binds or a promoter sequence comprising said SRE.

- 35. A method of identifying a potential insecticidal compound comprising:
 - (i) assaying the binding directly or indirectly of the recombinant or isolated polypeptide according to claim 30 to a steroid response element (SRE) to which said polypeptide binds, in the presence of a candidate compound; and
 - (ii) assaying the binding directly or indirectly of the recombinant or isolated polypeptide according to claim 30 to a steroid response element (SRE) to which said polypeptide binds, in the absence of said candidate compound; and
 - (ii) comparing the binding assayed at (i) and (ii), wherein a difference in the level of binding indicates that the candidate compound possesses potential insecticidal activity.
- 36. A method of identifying a candidate insecticidally-active agent comprising the steps of:

- a) expressing an EcR polypeptide of an insect steroid receptor or a fragment thereof which includes the ligand-binding region, optionally in association with an EcR partner protein (USP polypeptide) of an insect steroid receptor or ligand binding domain thereof, optionally in association with an insect steroid or analogue thereof so as to form a complex;
- b) purifying or precipitating the complex;
- determining the three-dimensional structure of the ligand binding domain of the complex; and
- d) identifying compounds which bind to or associate with the three-dimensional structure of the ligand binding domain, wherein said compounds represent candidate insecticidally-active agents.
- 37. A method of identifying a candidate insecticidally-active agent comprising the steps of:
 - a) expressing a USP polypeptide of a juvenile hormone receptor or a fragment thereof which includes the ligand-binding region, optionally in association with an EcR polypeptide of an insect steroid receptor or ligand binding domain thereof, and optionally in association with an insect steroid or analogue thereof, so as to form a complex;
 - b) purifying or precipitating the complex;
 - c) determining the three-dimensional structure of the ligand binding domain of the complex; and
 - d) identifying compounds which bind to or associate with the three-dimensional structure of the ligand binding domain, wherein said compounds represent candidate insecticidally-active agents.
- 38. A synthetic compound which interacts with the three dimensional structure of a polypeptide or protein selected from the list comprising:
 - (i) an EcR polypeptide of a steroid receptor or a fragment or bioactive derivative thereof:
 - (ii) an EcR partner protein (USP polypeptide) of a steroid receptor or a fragment or bioactive derivative thereof;

- 80 -

- (iii) a USP polypeptide of a juvenile hormone receptor or a fragment or bioactive derivative thereof; and
- (iv) a functional receptor or protein complex formed by association of (i) and (ii), wherein said compound is capable of binding to said polypeptide or protein to agonise or antagonise the binding activity or bioactivity thereof.
- 39. A method of identifying a synthetic compound for insecticidal activity comprising contacting the recombinant or isolated polypeptide according to claim 30 with said compound for a time and under conditions sufficient for binding to occur and detecting said binding using a detection means, wherein the occurrence of binding is indicative of potential insecticidal activity of the compound.

- 81 -

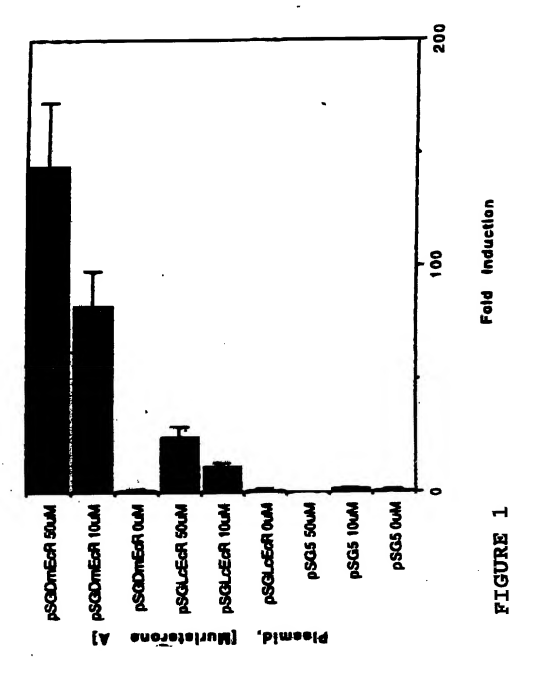
ABSTRACT

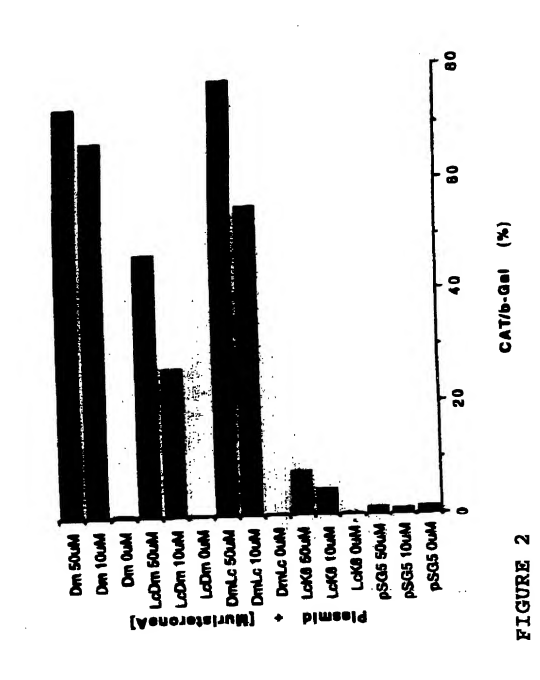
The present invention provides isolated nucleic acid molecules encoding polypeptides comprising functional steroid hormone and juvenile hormone receptors, in particular isolated nucleic acid molecules which encode polypeptides comprising the *Lucilia cuprina* and *Myzus persicae* ecdysone receptors and juvenile hormone receptors. The present invention further provides functional recombinant steroid and juvenile hormone receptors and recombinant polypeptide subunits thereof and derivatives and analogues thereof. The present invention further provides screening systems and methods of identifying insecticidally-active agents which are capable of agonising or antagonising insect receptor function, or alternatively or in addition, which modify the affinity of said receptors for their cellular stimuli (eg. insect steroids or juvenile hormones) or analogues thereof, or alternatively or in addition, which act as insecticides by viture of their ability to agonise or antagonise the activity of insect hormones.

15

10

5





3/4

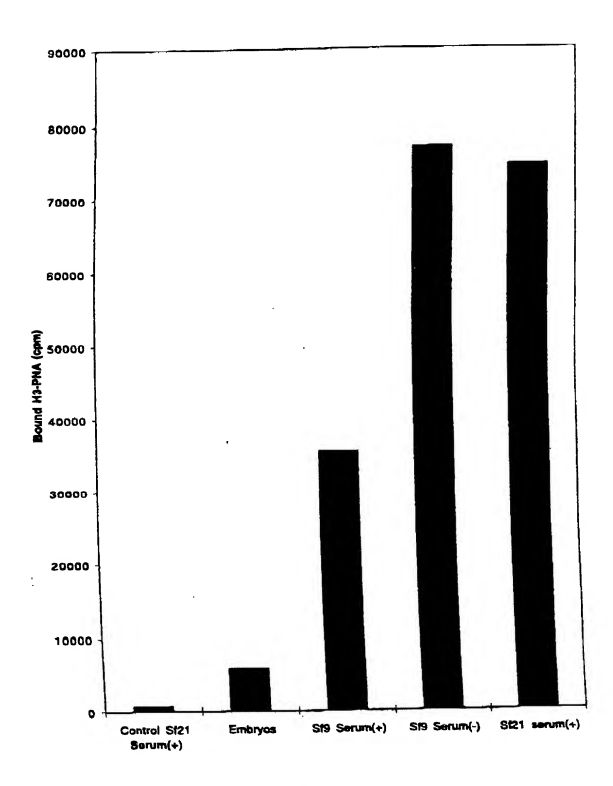


FIGURE 3

4/4

3H-PonA binding by Myzus in vitro translation products

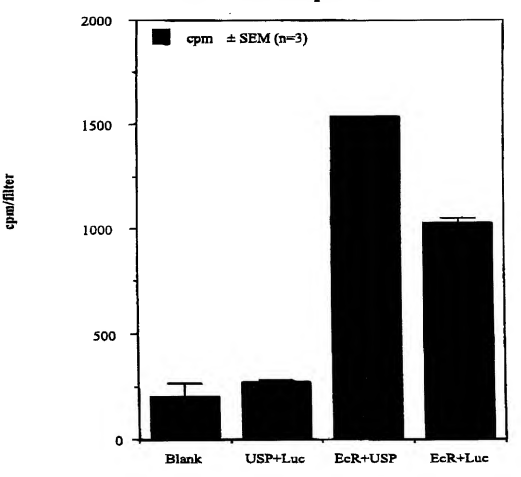


FIGURE 4

-1-

SEQUENCE LISTING

<110> COMMONWEALTH SCIENTIFIC AND INDUSTRIAL RESEARCH ORGANISATION <120> NOVEL GENETIC SEQUENCES ENCODING STEROID AND JUVENILE HORMONE 5 RECEPTOR POLYPEPTIDES AND INSECTICIDAL MODALITIES THEREFOR II <130> p:\oper\mro\ecdysone.cip <140> US continuation-in-part of PCT/AU99/00033 <141> 1999-07-02 10 <150> PCT/AU/00033 <151> 1999-01-15 <150> AU PP1536 <151> 1998-01-15 15 <160> 20 <170> PatentIn Ver. 2.0 <21,0> 1 20 <211> 2274 <212> DNA <213> Lucilia cuprina <220> 25 <221> CDS <222> (1)..(2271)

<400> 1

atg atg aaa cga cgt tgg tet aat aat ggc ggt ttt gcc gct tta aaa 48 Met Met Lys Arg Arg Trp Ser Asn Asn Gly Gly Phe Ala Ala Leu Lys

1

30

<

1,0

15

20

P:\OPER\MRO\ECDYSONE.CIP - 1/7/99

-2-

atg	tta	gaa	gaa	tcc	tac	tca	gaa	gta	acc	taa	taa	cca	aat	ggt	ctg	96
Met	Leu	Glu	Glu	Ser	Ser	Ser	Glu	Val	Thr	Ser	Ser	Ser	Asn	Gly	Leu	
			20					25					30			

- 5 gtd ttg tca tcg gat ata aat atg tca cct tcc tcg ttg gat tca ccc 144

 Val Leu Ser Ser Asp Ile Asn Met Ser Pro Ser Ser Leu Asp Ser Pro

 35 40 45
- gtt tat ggc gat cag gaa atg tgg ctg tgt aac gat tca gct tca tat 192

 10 Val Tyr Gly Asp Glm Glu Met Trp Leu Cys Asn Asp Ser Ala Ser Tyr

 50 55 60
 - Asn Asn Ser His Gln His Ser Val Ile Thr Ser Leu Gln Gly Cys Thr

 65 70 75 80
 - toa toa tig cog goo caa aca acc att ata cot cig toa got tia coc 288
 Ser Ser Leu Pro Ala Gln Thr Thr Ile Ile Pro Leu Ser Ala Leu Pro
 85 90 95

aat too aat aat goo too org aat aat caa aat caa aat tar caa aat 336
Asn Ser Asn Asn Ala Ser Leu Asn Asn Gln Asn Gln Asn Tyr Gln Asn
100 105 110

- 25 ggt aat tee atg aat aca aat tta teg gtt aac aca aat aac agt gtt 384
 Gly Asn Ser Met Asn Thr Asn Leu Ser Val Asn Thr Asn Asn Ser Val
 115
 120
 125
- gga gga ggt ggt ggt ggt ggt gta ccc ggt atg act tca ctc aat 432

 Gly Gly Gly Gly Gly Gly Gly Gly Val Pro Gly Met Thr Ser Leu Asn

 130

 130

 140

P;\OPER\MXO\ECDYSONE.CIF - 1/7/99

- 3 -

ggt	ctg	ggt	ggt	ggt	ggt	ggc	agt	ċąą	gtg	aat	aat	cac	aat	cac	age	480
Gly	Leu	Gly	Gly	Gly	Gly	Gly	Ser	Gln	Val	Asn	Asn	His	Asn	His	Ser	
145					150					155					160	
cac	aat	cat	tta	cac	cac	aac	agc	aa¢	agt	aat	cac	agt	aat	agc	agt	528
His	Asn	His	Leu	His	His	Asn	Ser	Asn	Ser	Asn	His	Ser	Asn	Ser	Ser	
				165					170					175		
teç	cac	cac	aca	aat	영영우	cac	atg	gg t	att	ggc	ggc	ggt	ggt	ggt	ggc	576
Ser	His	His	Thr	Asn	Gly	His	Met	Gly	Ile	Gly	Gly	Gly	Gly	Gly	Gly	
			180					185					190			
tta	ţcg	gtc	aat	att	aat	ggt	ccc	aat	atc	gtt	agc	aat	gcc	caa	cag	624
Leu	Ser	Val	Аsп	Ile	Asn	Gly	Pro	Asn	Ile	val	Ser	Asn	Ala	Gln	Gln	
		195					200					205				
tta	aac	teg	tta	cag	gcc	tca	caa	aat	gģc	caa	gtt	att	cat	gce	aat	672
Leu	Asn	Ser	Leu	Gln	Ala	Ser	Gln	Asn	Gly	Gln	Val	Ile	His	Ala	Asn	
	210					215			-		220					
att	gge	att	cac	a o rt.	atc	ate	aat	aa#	ccá	tta	aat	cat	ċ at	ca c	cat	720
		Ile														,,,
225			*****	5 41	230		5 01	ASII	GLY	235	USII	HIS	****	HID		
220					∪ دید					233					240	
		_			_	-	_	_					•		gaa	768
HIS	HIB	Mec	Asn		Ser	ser	Met	Met		His	Thr	Pro	Arg		Glu	
				245					250					255		
									_	-				_	agc	816
Ser	Ala	Asn	Ser	Ile	Ser	Ser	СĴУ	Arg	Asp	Asp	Leu	Ser	Pro	Ser	Ser	

10

15

20

25

370

P:\QPER\MRQ\ECDYSONE,CIP - 1/7/99

-4-

agt	ctt	aat	ggc	ttc	tca	aca	agc	gat	gct	agt	gat	gtt	aag	aaa	atc	864
Ser	Leu	Asn	Gly	Phe	Ser	Thr	Ser	Asp	Ala	ser	Asp	Val	Lys	Lys	Ile	
		275					280					285				
aaa	aaa	33¢	cct	gcg	ccc	cgt	tta	caa	aaa	gaa	ctg	tgt	ctg	gtg	tgt	912
ГÀз	Lys	Gly	Pro	Ala	Pro	Arg	Leu	Gln	Glu	Glu	Leu	Сла	Leu	Val	cha	
	290					295					300					
ggt	gat	cgg	gcg	tac	ggt	tat	cat	tat	aac	gca	ctc	acc	tgt	gaa	99c	960
Gly	Asp	Arg	Ala	Ser	Gly	Tyr	His	Tyr	Asn	Ala	Leu	Thr	Сув	Glu	Gly	
305					310					315					320	
tgt	aag	3 99	ttc	ttt	cga	cgg	agt	gtt	acc	aaa	aat	ತಿငಡಿ	gtg	tat	tgt	1008
Cys	Lys	Gly	Phe	Phe	Arg	Arg	Ser	Val	Thr	Lys	Asn	Ala	Val	Tyr	Суз	
				325					330					335		
tgt	aaa	ttt	ggt	cat	300	tgc	gaa	atg	gac	atg	tat	atg	cds	cgt	aaa	1056
Cys	Lys	Phe	Gly	His	Ala	СЛа	Glu	Met	Asp	Met	тут	Met	Arg	Arg	Lys	
			340					345					350			
tgt	cag	gaa	tgt	ಕ್ರಡ	ctg	aaa	aaa	tgt	ttg	gct	gtg	ggc	atg	cgg	ceg	1104
Cys	Gln	Glu	Суз	Arg	Leu	Lys	Lys	CAa	Leu	Ala	Val	Gly	Met	Arg	Pro	
		355					360					365				
ass des	tgt	gtg	gtg	CCC	gaa	aac	cag	tgt	gca	atg	aaa	cga	cgc	gaa	aag	1152

Glu Cys Val Val Pro Glu Asn Gln Cys Ala Met Lys Arg Arg Glu Lys

380

375

10

15

20

25

30

P:\OPER\MRO\ECDYSONE.CIP - 1/1/99

- 5 -

aaa	gca	caa	aaa	gag	aag	gat	aaa	ata	cag	acc	agt	gtg	tgt	gca	acg	1200
Lys	Ala	Gln	ГÀв	Glu	Lys	Asp	Lys	Ile	Gln	Thr	Ser	val	Cys	Ala	Thr	
385					390					395					400	
gaa	act	aaa	aag	gaa	ata	ctc	gat	tta	atg	aca	tgt	gaa	ccg	cca	tca	1248
Glu	Ile	Lys	Lys	Glu	Ile	Leu	Asp	Leu	Met	Thr	Cys	Glu	Pro	Pro	Ser	
				405					410					415		
cat	cca	acg	tgţ	ccg	ctg	tta	cct	gaa	gac	att	ttg	gct	aaa	tgt	caa	1296
His	Pro	Thr	Сув	Pro	Leu	Leu	Pro	Glu	Asp	Ile	Leu	Ala	Lys	Суз	Gln	
			420					425					430			
gct	egt	aat	ata	cct	cct	tta	tcg	tac	aat	caa	ttg	àcə	gtt	ata	tat	1344
Ala	Arg	Asn	Ile	Pro	Pro	Leu	Ser	Тут	Asn	Gln	Leu	Ala	Val	Ile	Tyr	
		435					440					445				
aaa	tta	aca	tgg	tat	caa	gat	ggc	tac	gaa	cag	cca	tcc	gag	gaa	gat	1392
Lys	Leu	Ile	Trp	Tyr	Gln	Asp	Gly	Tyr	Glu	Gln	Pro	Ser	Glu	Glu	Asp	
	450					455					460					
ctc	aaa	cgt	ata	atg	agt	tça	ccc	gat	gaa	aat	gaa	agt	caa	cac	gat	1440
Leu	Lys	Arg	Ile	Met	Ser	Ser	Pro	Asp	Glu	Asn	Glu	Ser	Gln	His	Asp	
465					470					475					480	
gca	tca	ttt	cgt	cat	ata	aca	gaa	atc	act	ata	cta	aca	gta	Caa	tta	1488
Ala	Ser	Phe	Arg	His	Ile	Thr	Glu	Ile	Thr	Ile	Leu	Thr	Val	Gln	Leu	
				485					490					495		
att	gtg	gaa	ttt	gcc	aag	åã₽	ttg	cca	ācā	ttt	acc	aaa	ata	cca	caa	1536
Ile	Val	Glu	Phe	Ala	Lys	Gly	Leu	Pro	Ala	Phe	Thr	ГÀЗ	Ile	Pro	Gln	
			500					505					510			

P:\OPER\MRO\ECDYSONE.CIP - 1/7/99

-6-

gag	gat	caa	ata	aca	cta	tta	aag	gcc	tgc	tca	tca	gaa	gtt	atg	atg	1584
Glu	Asp	Gln	Ile	Thr	Геп	Leu	Lys	Ala	Cys	Ser	Ser	Glu	Val	Met	Met.	
		515					520					525				
ttg	cga	atg	gca	cga	cgt	tac	gat	cac	aat	tca	gat	tog	ata	ttc	ttt	1632
Leu	Arg	Met	Ala	Arg	Arg	тут	Asp	His	Asn	Ser	Авр	Ser	Ile	Phe	Phe	
	530					535					540					
cc a	at a	nat c	ga t	cg t	at a	යේ ර	gt g	gac t	et t	at a	aa a	itg ç	get g	iãc s	atg	1680
Ala	Asn	Asn	Arg	Ser	Tyr	Thr	Arg	Asp	Ser	Tyr	Lys	Met	Ala	Gly	Met	
545					550					555					560	
gct	gat	aat	att	gag	gaț	ctg	ctg	cat	ttc	tgt	cga	caa	atg	tac	teg	1728
Ala	Asp	Asn	Ile	Glu	Asp	Leu	Leu	His	Phe	Сув	Arg	Gln	Met	Tyr	Ser	
				565					570					575		
atg	aaa	gtg	gac	aat	gtc	gaa	tat	gct	cta	ctc	act	gee	att	gtg	atc	1776
Met	Lys	v al	Asp	Asn	Val	Glu	Tyr	Ala	Leu	Leu	Thr	Ala	Ile	Val	Ile	
			580					585					590			
ttt	tcc	gat	¢gg	ccq	ggt	ete	gaa	gaa	acc	qaa	cta	ata	qaa	aca	ata	1824
			Arg													
		595	5	,,,,	1		600					605			-10	
		525					000					ران				
caa	agt	tac	tac	atc	gat	aca	ctc	අදිය	att	tac	ata	ctt	aat	cgc	cat	1872
Gln	Ser	Туг	Tyr	Ile	Asp	Thr	Leu	Arg	Ile	Tyr	Ile	Leu	Asn	Arg	His	
	610					615					620					
tgc	ggc	gat	cc¢	atg	agt	ctc	gta	ttc	ttt	gcc	aag	ctt	ctg	tca	att	1920
Cys	Gly	Asp	Pro	Met	Ser	Leu	Val	Phe	Phe	Ala	Lys	Leu	Leu	Şer	Ile	

10

15

20

25

. 30

P:\OPER\MRO\ECDYSONE.CIP - 1/7/99

-7-

cta	acc	gaa	ctg	cgt	acg	ttg	ggc	aat	caa	aat	gcc	gaa	atg	tgt	ttc	1968
Leu	Thr	Glu	Leu	Arg	Thr	Leu	Gly	Asn	Gln	Asn	Ala	Glu	Met	Cys	Phe	
				645					650					655		
tcg	ttg	aaa	ttg	aaa	aat	cgc	aaa	ctg	cca	aaa	ttc	ata	gaa	gag	atc	2016
Ser	Leu	Lys	Leu	Lув	Asn	Arg	ГÀЗ	Leu	Pro	Lys	Phe	Leu	Glu	Glu	Ile	
			660					665					670			
Fgg	gat	gta	cat	gec	att	cca	ccc	tca	gtg	cag	tca	cac	ata	cag	gct	2064
Trp	Asp	Val	His	Ala	Ile	Pro	Pro	Ser	Val	Gln	Ser	His	Ile	Gln	Ala	
		67 5					680					685				
acc	cag	àcā	gaa	aag	àcc	âcc	cag	gaa	gct	cag	gca	aca	aca	tcg	gcc	2112
Thr	Gln	Ala	Glu	Lys	Ala	Ala	Gln	Glu	Ala	Gln	Ala	Thr	Thr	Ser	Ala	
	690					695					700					
att	tca	gca	gcc	gac	ac¢	tca	tct	tcc	taa	ata	aat	acc	tcg	atg	acs	2160
Ile	Ser	Ala	Ala	Ala	Thr	Ser	Ser	Ser	Ser	Ile	Asn	Thr	Ser	Met	Ala	
705					710					715					720	
aca	tca	tcc	tca	tca	teg	tta	tog	೮೦೩	tcg	gcg	gaa	tda	aca	ccc	aat	2208
Thr	Ser	Ser	Ser	Ser	Ser	Leu	Ser	Pro	Ser	Ala	Ala	Ser	Thr	Pro	Asn	
				725					730					735		
ggt	ggt	gce	gtc	gat	tat	gtt	ggc	acc	gat	atg	agt	atg	agt	tta	gta	2256
Gly	Gly	Ala	Val	Asp	Tyr	Val	Gly	Thr	Asp	Met	Ser	Met	Ser	Leu	Val	
			740					745					750			
caa	tcg	gat	aat	gca	tag	•										2274
Ģln	Ser	Asp	Asn	Ala												

P:\OPER\MRO\ECDYSONE,CIP - 1/1/99

-8-

<210> 2

<211> 757

5 <212> PRT

<213> Lucilia cuprina

<400> 2

Met Met Lys Arg Arg Trp Ser Asn Asn Gly Gly Phe Ala Ala Leu Lys

10 1 5 10 15

Met Leu Glu Ser Ser Ser Glu Val Thr Ser Ser Ser Asn Gly Leu
20 25 30

Val Leu Ser Ser Asp Ile Asn Met Ser Pro Ser Ser Leu Asp Ser Pro4045

Val Tyr Gly Asp Gln Glu Met Trp Leu Cys Asn Asp Ser Ala Ser Tyr
50 55 60

Asn Asn Ser His Gln His Ser Val Ile Thr Ser Leu Gln Gly Cys Thr

65 70 75 80

Ser Ser Leu Pro Ala Gln Thr Thr Ile Ile Pro Leu Ser Ala Leu Pro

25

85

90

95

Asn Ser Asn Asn Ala Ser Leu Asn Asn Gln Asn Gln Asn Tyr Gln Asn 100 105 110

30 Gly Asn Ser Met Asn Thr Asn Leu Ser Val Asn Thr Asn Asn Ser Val

-9-

Gly Gly Gly Gly Gly Gly Gly Val Pro Gly Met Thr Ser Leu Asn 130 135 140

Gly Leu Gly Gly Gly Gly Ser Gln Val Asn Asn His Asn His Ser

5 145 150 155 160

His Asn His Leu His His Asn Ser Asn Ser Asn His Ser Asn Ser Ser 165 170 175

10 Ser His His Thr Asn Gly His Met Gly Ile Gly Gly Gly Gly Gly Gly 180 185 190

Leu Ser Val Asn Ile Asn Gly Pro Asn Ile Val Ser Asn Ala Gln Gln
195 200 205

15

Leu Asn Ser Leu Gln Ala Ser Gln Asn Gly Gln Val Ile His Ala Asn 210 215 220

Ile Gly Ile His Ser Ile Ile Ser Asn Gly Leu Asn His His His His 20 225 230 235 240

His His Met Asn Asn Ser Ser Met Met His His Thr Pro Arg Ser Glu
245 250 255

25 Ser Ala Asn Ser Ile Ser Ser Gly Arg Asp Asp Leu Ser Pro Ser Ser
260 265 270

Ser Leu Asn Gly Phe Ser Thr Ser Asp Ala Ser Asp Val Lys Lys Ile 275 280 285

30

Lys Lys Gly Pro Ala Pro Arg Leu Gln Glu Glu Leu Cys Leu Val Cys

- 10 -

290

295

300

Gly Asp Arg Ala Ser Gly Tyr His Tyr Asn Ala Leu Thr Cys Glu Gly
305 310 315 320

5

Cys Lys Gly Phe Phe Arg Arg Ser Val Thr Lys Asn Ala Val Tyr Cys
325 330 335

Cys Lys Phe Gly His Ala Cys Glu Met Asp Met Tyr Met Arg Arg Lys

10 340 345 350

Cys Gln Glu Cys Arg Leu Lys Lys Cys Leu Ala Val Gly Met Arg Pro 355 360 365

15 Glu Cys Val Val Pro Glu Asn Gln Cys Ala Met Lys Arg Arg Glu Lys370 375 380

Lys Ala Gln Lys Glu Lys Asp Lys Ile Gln Thr Ser Val Cys Ala Thr
385 390 395 400

20

Glu Ile Lys Lys Glu Ile Leu Asp Leu Met Thr Cys Glu Pro Pro Ser 405 410 415

His Pro Thr Cys Pro Leu Leu Pro Glu Asp Ile Leu Ala Lys Cys Gln
25 420 425 430

Ala Arg Asn Ile Fro Pro Leu Ser Tyr Asn Gln Leu Ala Val Ile Tyr
435 440 445

30 Lys Leu Ile Trp Tyr Gln Asp Gly Tyr Glu Gln Pro Ser Glu Glu Asp
450 460

25

P:\OPER\MRO\ECDYSONE.CIP - 1/7/99

515

- 11 -

Leu Lys Arg Ile Met Ser Ser Pro Asp Glu Asn Glu Ser Gln His Asp 465 470 475 480

Ala Ser Phe Arg His Ile Thr Glu Ile Thr Ile Leu Thr Val Gln Leu

5 495 495

Ile Val Glu Phe Ala Lys Gly Lou Pro Ala Phe Thr Lys Ile Pro Gln
500 505 510

Glu Asp Gln Ile Thr Leu Leu Lys Ala Cys Ser Ser Glu Val Met Met

Leu Arg Met Ala Arg Arg Tyr Asp His Asn Ser Asp Ser Ile Phe Phe

15 530 535 540

520

525

Ala Asn Asn Arg Ser Tyr Thr Arg Asp Ser Tyr Lys Met Ala Gly Met
545 550 555 560

20 Ala Asp Asn Ile Glu Asp Leu Leu His Phe Cys Arg Gln Met Tyr Ser
565 570 575

Met Lys Val Asp Asn Val Glu Tyr Ala Leu Leu Thr Ala Ile Val Ile . 580 585 590

Phe Ser Asp Arg Pro Gly Leu Glu Glu Ala Glu Leu Val Glu Ala Ile
595 600 605

Gln Ser Tyr Tyr Ile Asp Thr Leu Arg Ile Tyr Ile Leu Asn Arg His

620

- 12 -

Cys Gly Asp Pro Met Ser Leu Val Phe Phe Ala Lys Leu Leu Ser Ile
625 630 635 640

Leu Thr Glu Leu Arg Thr Leu Gly Asn'Gln Asn Ala Glu Met Cys Phe
645 650 655

Ser Leu Lys Leu Lys Asn Arg Lys Leu Pro Lys Phe Leu Glu Glu Ile
660 665 670

10 Trp Asp Val His Ala Ile Pro Pro Ser Val Gln Ser His Ile Gln Ala 675 680 685

Thr Gln Ala Glu Lys Ala Ala Gln Glu Ala Gln Ala Thr Thr Ser Ala 690 695 700

15

5

Ile Ser Ala Ala Ala Thr Ser Ser Ser Ser Ile Asn Thr Ser Met Ala
705 710 710 715 720

Thr Ser Ser Ser Ser Leu Ser Pro Ser Ala Ala Ser Thr Pro Asn 20 725 730 735

Gly Gly Ala Val Asp Tyr Val Gly Thr Asp Met Ser Met Ser Leu Val
740 745 750

25 Gln Ser Asp Asn Ala

755

<210> 3

<211> 1401

30 <212> DNA

<213> Lucillia cuprina

80

P:\OPER\MRO\ECDYSONE,CIP - 1/7/99

- 13 -

<220>

<221> CDS

<222> (1)..(1401)

5 <400> 1 atg gat aac ggc gag daa gat gct ggg tto cga ttg gca ccg atg tct Met Asp Asn Gly Glu Gln Asp Ala Gly Phe Arg Leu Ala Pro Met Ser 10 15 15 ccg cag gag ata aag cca gac att tca cta ctc aat gaa aat aat acg Pro Gln Glu Ile Lys Pro Asp Ile Ser Leu Leu Asn Glu Asn Asn Thr 20 agt agt tat tog coc aaa cot gga agt cot aat coa tit gec atc gga 144 25 Ser Ser Tyr Ser Pro Lys Pro Gly Ser Pro Asn Fro Phe Ala Ile Gly 35 40 30 ttg cag gca ata aat gca gtc gct gcc gcg aat gcc aat aac caa aat 192 Leu Gln Ala Ile Asn Ala Val Ala Ala Ala Asn Ala Asn Asn Gln Asn 35 50 caa atg ttg caa act acg cca cca caa cag cag cag tat cca cca aat 240 40 Gln Met Leu Gln Thr Thr Pro Pro Gln Gln Gln Tyr Pro Pro Asn 70 75 45 cae eec ett agt ggt teg aaa eac ttg tgt tee att tgt gga gae ege His Pro Leu Ser Gly Ser Lys His Leu Cys Ser Ile Cys Gly Asp Arg 50 в5 95 55 gcc agt gga aaa cat tat ggg gtc tac agt tgt gag ggt tgt aaa ggg 336 Ala Ser Gly Lys His Tyr Gly Val Tyr Ser Cys Glu Gly Cys Lys Gly 100 105 60 tto tto aaa ogt acc gta ogc aag gao ttg aca tat get tgt ogt gag 384 65 Phe Phe Lys Arg Thr Val Arg Lys Asp Leu Thr Tyr Ala Cys Arg Glu 115 120 70 gad aga aat tgc att ata gat aaa cga daa aga aat dgt tgc dag tat 432 Asp Arg Asn Cys Ile Ile Asp Lys Arg Gln Arg Asn Arg Cys Gln Tyr 75 135 tgt cgt tat caa aag tgt tta gct tgt ggc atg aaa cgc gaa gcg gtc

Cys Arg Tyr Gln Lys Cys Leu Ala Cys Gly Met Lys Arg Glu Ala Val

- 14 -

	145					150					155					160	
5	caa	gag	gaa	cga	caa	cgt	ggt	act	cgt	gct	gat	aac	gct	aga	got	gct	528
	Gln	Glu	Glu	Arg	Gln	Arg	Gly	Thr	Arg	Ala	Ala	Asn	Ala	Arg	Ala	Ala	
10					165					170				_	175		
10																	
	33¢	gct	ggc	ggt	99¢	gga	gga	ggt	ggt	ggt	333	gta	agc	aat	gtg	gtt	576
15	Gly	Ala	Gly	Gly	Gly	Gly	Gly	Gly	Gly	Gly	Gly	val	Ser	Asn	Val	Val	
				180					185					190			
20																	
					gaa										_		624
	Gly	Ala	Ģly	Gly	Glu	Asp	Phe	Г À з	Pro	Ser	Ser	Ser	Leu	Arg	Asp	Leu	
25			195					200					205				
30					atc												672
	Thr		Glu	Arg	Ile	Ile		Ala	Glu	Gln	Lys		Glu	Ser	Leu	Ser	
35		210					215					220					
JĢ		~~+		ar							_						
					ttg												720
40	225	veb	WOII	VAL	Leu	230	Fne	ren	Arg	AST		Asn	ASN	Ser	Met		
	227					230					235					240	
45	caa	cac	gac	tac	aaa	dac	aca	ort.a	tot	cat		tac	Cac	ato	~++	226	768
					Lys												100
			-		245	,				250		-,-	- LI		255	rion.	
50										220							
	aaa	Çaa	cta	tac	caa	atg	gtt	gaa	tat	gca	cgt	cqa	aca	cca	cat	ttt	816
55					Gln												
				260					265		_			270			
60																	
00	aca	cat	ttg	çag	cgt	ಡಿತಡ	gat	cag	ata	cta	ttg	tta	aag	gct	3 3c	tgg	864
	Thr	His	Lęц	Gln	Azg	Glu	Asp	Gln	Ile	Leu	Leu	Leu	Lys	Ala	Gly	Trp	
<i>6</i> 5			275					280					285				
70					att												912
	Asn	Glu	Leu	Leu	Ile	Ala	Asn	Val	Ala	Trp	Ċys	Ser	Ile	Glu	Ser	Leu	
		290					295					300					
75																	
					gcc												960
80		Ala	Glu	Tyr	Ala		Pro	Gly	Thr	Val	His	Asp	Gly	Ser	Phe	Gly	
	305					310					315					320	
25			a														
85					gtg												100\$
	Arg	Arg	ser	FLO	Val	Arg	Gln	Pro	GLn		Leu	Phe	Leu	Asn		Asn	
90					325					330					335		

- 15 -

		_			ege		_	· .		_	_		-	-			1056
5	rne	ser	ıyı	340	Arg	ASI	ser	Ala	345	пÀв	WTS	ASII	AST	350	ser	TTG	
10		-	Arg		ctc Leu	_		Leu	•			_	Lys				1104
15	atc	gat	355 cac	tea	gag	tta	tea	360 tat	cta	ຄອຕ	dr.a.	atc	365 ara	et c	t t c	== !	1152
20					Glu												1172
25					ggt					-				_	_	_	1200
30	385					390					395					400	
35	gaa Glu				gcc Ala 405												1248
40					cgc											-	1296
45	GIY	Asp	Asp	420	Arg	РДе	Ala	Gln	425	Leu	Leu	Arg	Leu	Pro 430	Ala	Leu	
50	cgt Arg				ctc Leu										-		1344
55	252	~~-															
60					Ala							_			_		1392
65	cct Pro		_														1401
70	465																
75	<210)> 4 .> 46	57														
80		!> PI !> L\		lia (cupr:	ina											
85	<400				.				-								
90	Met 1	Asp	Asn	Gly	Glu 5	Gln	Asp	Ala	Gly	Phe 10	Arg	Leu	Ala	Pro	Met 15	Ser	

- 16 -

	Pro	Gln	Glu	Ile 20	Lys	Pro	Asp	Ile	Se <i>r</i> 25	Leu	Leu	Asn	Glu	Asn 30	Asn	Thr
5	Ser	Ser	Tyr	Ser	Pro	Lys	Pro	Gly	Ser	Pro	Asn	Pro	Phe	Ala	Ile	Gly
10			35			-		40					45			1
15	Leu	Gln 50	Ala	Ile	Asn	Ala	Val 55	Ala	Ala	Ala	Aşn	Ala 60	Asn	Asn	Gln	Asn
20	Gln 65	Met	Leų	Gln	Thr	Th r 70	Pro	Pro	Gln	Gln	Gln 75	Gln	Тут	Pro	Pro	Asn 80
25	His	Pro	Leu	Ser	Gly 85	Ser	Lys	His	Leu	Су в 90	Ser	Ile	Сув	Gly	Asp 95	Arg
30 35	Ala	Ser	Gly	Lys 100	His	туг	Gly	Val	Тут 105	Ser	СХз	Glu	Gly	Cys 110	Lys	Gly
40	Phe	Phe	Lys 115	Arģ	Thr	Val	Arg	Lys 120	Asp	Leu	Thr	Tyr	Ala 125	Сув	Arg	Glu
45	Asp	Arg 130	naA	Суѕ	Ile	Ile	Asp 135	ŗåa	Arg	Gln	Arg	Asn 140	Arg	Cys	Gln	Tyr
50	Cys 145	Arg	Tyr	Gln	Lys	Cys 150	Геп	Ala	Сув	Gly	Met 155	Lys	Arg	Glu	Ala	Val 160
5 5	Gln	Glu	Glu	Arg	Gln 165	Arg	Gly	Thr	Arg	Ala 170	Ala	Asn	Ala	Arg	Ala 175	Ala
60 65	Gly	Ala	Gly	Gly 180	Gly	Gly	Gly	Gly	Gly 185	Gly	Gly	Val	Ser	Asn 190	Val	Val
70	Gly	Ala	Gly 195	Gly	Glu	Asp	Phe	L y s 200	Pro	Ser	Ser	Ser	Leu 205	Arg	Asp	Leu
75	Thr	Ile 210	Glu	Arg	Ile	Ile	Glu 215	Ala	Glu	Gln	Lys	Ala 220	Glu	Ser	Leu	Ser
80	Gly 225	Ąsp	Asn	Val	Leu	Pro 230	Phe	Leu	Arg	Val,	Gly 235	Asn	Asn	Ser	Met	Val 240
85	Gln	His	Asp	Tyr	Lys 245	Gly	Ala	Val	Ser	His 250	Leu	Сув	Gln	Met	Val 255	Asn
90	Lvs	Gln	Leu	ጥህታ	Gla	Met	va1	Glu	7	Z 7 =	725	71	πh	Dec	13 -	₽F -

260

- 17 -

				260					265					270		
5	Thr	His	Leu 275	Gln	Arg	Glu	Asp	Gln 280	Ile	Leu	Leu	Leu	Lys 285	Ala	Gly	Trp
10	Asn	Glu 290	Leu	Leu	Ile	Ala	Asn 295	Val	Ala	Trp	Cys	Ser 300	Ile	Glu	Ser	Leu
15	Ая р 305	Ala	Glu	Туг	Ala	Ser 310	Pro	Gly	Thr	Val	His 315	Asp	Gly	Ser	Phe	
20		A	Som	Dwa	V-1		C1	n	a) -	63 -		mk -	T	3	61 -	320
25	Atg	wrg	ser	FIC	Val 325	urg	GIN	FIO	GIII	330	ьец	Pne	Leu	Asn	335	Asn
30	Phe	Ser	Tyr	His 340	Arg	Asn	Sex	Ala	Ile 345	Lys	Ala	Asn	Val	Val 350	Ser	Ile
35	Phę	Asp	Arg 355	Ile	Leu	Ser	Glu	Leu 360	Ser	Ile	Lys	Met	Lys 365	Arg	Leu	Asn
40 45	Ile	Asp 370	Arg	Ser	Glu	Leu	Ser 375	Сув	Leu	ŗÅŝ	Ala	Ile 380	Ile	Leu	Phe	Asn
50	Pro 385	Asp	Ile	Arg	Gly	Leu 390	Lys	Cys	Arg	Ala	Asp 395	Val	Glu	Val	Сув	Arg 400
55	Glu	Ľys	Ile	Tyr	Ala 405	Суз	Leu	Asp	Glu	His 410	Сув	Arg	Thr	Glu	His 415	Pro
60	Gly	Авр	Asp	Gly 420	Arg	Phe	Ala	Gln	Leu 425	Leu	Ге́л	Arg	Leu	Pro 430	Ala	Leu
65	Arg	Ser	Ile 435	Ser	Leu	Lys	Cys	Leu 440	Asp	His	Leu	Phe	Phe 445	Phe	Arg	Leu
70	Īle	Gly 450	Glu	Arg	Ala	Leu	Glu 455	Glu	Гéй	Ile	Ala	Glu 460	Gln	Leu	Glu	Ala
75	Dro	T 16	Carc													
80	465	Ile	Ċλὰ													

- 18 -

<210> 5

<211> 585

<212> DNA

<213> Myzus persicae

5

<220>

<221> CDS

<222> (1)..(585)

10 <400> 5

gaa ttc ggc acg age gcc att gtt aat gga ttt atc cgc acc att agt 48
Glu Phe Gly Thr Ser Ala Ile Val Asn Gly Phe Ile Arg Thr Ile Ser

1 5 10 15

15 ttg atc ctt att ttt ctt ctt ctt ttt ctt tgg agg ttg ttg gec ttc 96

Leu Ile Leu Ile Phe Leu Leu Phe Leu Trp Arg Leu Leu Ala Phe

20 25 30

egg ttc ttg ttt ata tct gaa caa cca cct ccc gaa gag ctg tgc ctg 144

20 Arg Phe Leu Phe Ile Ser Glu Glu Pro Pro Pro Glu Glu Leu Cys Leu

35 40 45

gtg tgt gge gad ogg teg tee ggt tae eat tae aac get etc aca tgc 192

Val Cys Gly Asp Arg Ser Ser Gly Tyr His Tyr Asn Ala Leu Thr Cys

50 55 60

gaa gga tgc aag ggg ttc ttc cgg agg agc atc acc aag aac gcc gtg 240 Glu Gly Cys Lys Gly Phe Phe Arg Arg Ser Ile Thr Lys Asn Ala Val 65 70 75 80

30

25

tac cag tgc aag tac ggc aac aat tgc gaa atc gac atg tac atg agg 288

195

P.\OPER\MRO\ECDYSONE,C(P-1/7/99

- 19 -

	Tyr	Gln	Cys	Lys	Tyr	Gly	Asn	Asn	Сув	Glu	Ile	Asp	Met	Tyr	Met	Arg	
					85					90					95		
	cgg	aag	tgc	cag	gag	tgc	cââ	ctg	aaa	aaa	tg¢	ctg	acc	gtc	ggc	atg	336
5	Arg	Lys	Сув	Gln	Glu	Cys	Arg	Leu	Lys	Lys	Суз	Leu	Thr	Val	Gly	Met	
				100					105					110			
	g å å	cct	gaa	tgt	gtt	gta	cct	gaa	gtt	caa	tgc	gca	gta	222	aga	aag	384
	Arg	Pro	Glu	Сув	V al	val	Pro	Glu	Val	Gln	Cys	Ala	Val	ГÀЗ	Arg	Lys	
10			115					120					125				
	gag	aaa	aaa	gct	caa	cga	gaa	aaa	gat	aaa	cca	aat	tat	act	aca	gac	432
	Glu	Lys	Lys	Ala	Gln	Arg	Glu	Lys	Asp	Lys	Pro	Asn	Ser	Thr	Thr	Asp	
		130					135					140					
15																	
	att	tot	cct	gaa	ata	ata	aaa	ata	gaa	cct	aca	gag	atg	aag	att	gaa	480
	Ile	Ser	Pro	Glu	Ile	Ile	Lys	Ile	Glu	Pro	Thr	Glu	Met	Lys	Ile	Glu	
	145					150					155					160	
20	tgt	ggt	gaa	cca	atg	ata	atg	3 3 c	aca	CCE	atg	ccg	act	gta	cct	tac	528
	Cys	Gly	Glu	Pro	Met	Ile	Met	Gly	Thr	Pro	Met	Pro	Thr	Val	Pro	Tyr	
					165					170					175		
	gtg	aaa	cct	ttg	agt	tct	ctc	gtg	ccg	aat	tcg	ācs	cga	gta	a¢g	ggt	576
25	Val	Lys	Pro	Leu	Ser	Ser	Leu	Val	Pro	Asn	Ser	Ala	Arg	Val	Thr	Gly	
				180					185					190			
	tac	aaa	EEE														585
	Tyr	Lys	Phe														

- 20 -

<210> б

<211> 195

<212> PRT

5 <213> Myzus persicae

<400> 6

Glu Phe Gly Thr Ser Ala Ile Val Asn Gly Phe Ile Arg Thr Ile Ser

1 5 10 15

10

Leu Ile Leu Ile Phe Leu Leu Phe Leu Trp Arg Leu Leu Ala Phe

20 25 30

Arg Phe Leu Phe Ile Ser Glu Gln Pro Pro Pro Glu Glu Leu Cys Leu

15 35 40 45

Val Cys Gly Asp Arg Ser Ser Gly Tyr His Tyr Asn Ala Leu Thr Cys

50 55 60

20 Glu Gly Cys Lys Gly Phe Phe Arg Arg Ser Ile Thr Lys Asn Ala Val

65 70 75 80

Tyr Gln Cys Lys Tyr Gly Asn Asn Cys Glu Ile Asp Met Tyr Met Arg

95 90 95

25

Arg Lys Cys Gln Glu Cys Arg Leu Lys Lys Cys Leu Thr Val Gly Met

100 105 110

Arg Pro Glu Cys Val Val Pro Glu Val Gln Cys Ala Val Lys Arg Lys

30 115 120 125

-21 -

Glu Lys Lys Ala Gln Arg Glu Lys Asp Lys Pro Asn Ser Thr Thr Asp 130 135 140

Ile Ser Pro Glu Ile Ile Lys Ile Glu Pro Thr Glu Met Lys Ile Glu

145 150 155 160

Cys Gly Glu Pro Met Ile Met Gly Thr Pro Met Pro Thr Val Pro Tyr

165 170 175

10 Val Lys Pro Leu Ser Ser Leu Val Pro Asn Ser Ala Arg Val Thr Gly
180 185 190

Tyr Lys Phe

195

15

5

<210> 7 <211> 208 <212> DNA <213> Myzus persicae

20

25

<400> 7
catgootgoa ggtogactot agaggatoco otogtocggt taccattaca acgcactoac 60

ctgtgaaggc tgtaagggtt tctttcgacg gagtgttacc aaaaatgcgg tgtattgttg 120
taaatttggt catgcctgcg aaatggacat gtatatgcga cgtaaatgtc aggaatgtag 180
gctgaaaaaa tgtttggctg tgggcatg 208

30

35

<210> 8 <211> 436 <212> DNA <213> Myzus persicae

<400> 8 catgeggeeg gaatgtgtgg tgcccgaaaa ccagtgtgca atgaaacgac gcgaaaagaa 60

- 22 -

	agcacaaaaa gagaaggata aaatacagac cagtgtgtgt gcaacgg	aaa ttaaaaagga 120
	aatactcgat ttaatgacat gtgaaccgcc atcacatcca acgtgtc	cgc tgttacctga 180
5	agacattitg getaaatgte aagetegtaa tataceteet ttategt	aca atcaattggc 240
	agttatatat aaattaatat ggtatcaaga tggctacgaa cagccat	ccg aggaagatet 300
10	caaacgtata atgagtteae eegatgaaaa tgaaagteaa caegatg	cat catttegtca 360
,,,	tataacagaa atcactatac taacagtaca attaattgtt gaatgtg	cca aaggtctagg 420
	gtaccgagct cgaatt	436
15		
	<210> 9	
20		
20	<211> 1797	
	<212> DNA	
25	<213> Myzus persicae	
	<220>	
30	<221> CDS	
	<222> (1),.(1797)	
A-		
35	<400> 9	
	atg atg gac cag asa tgt gac gtc ggc ggt ggt ggt gtc	get get gee 48
40	Met Met Asp Gln Lys Cys Asp Val Gly Gly Gly Val	
	1 5 10	15
	1 3	43
45		
	gee gee ggt are ggt gge ggt gte gge gge ete atg	
	Ala Ala Gly Ile Gly Gly Gly Val Gly Gly Leu Met	_
50	20 25	30
55	egt gge egt gge gge add gag gte ate ate aaa eee egt	agt cot gcc 144
55	Arg Gly Arg Gly Gly Thr Glu Val Ile Ile Lys Pro Arg	g Ser Pro Ala
	35 40 45	5
60		
	gtg gtg cag gtg gcc acc ggt ggc agt tac cac ggc etg	g eeg geg gee 192
	Val Val Gln Val Ala Thr Gly Gly Ser Tyr His Gly Let	
65	· · ·	1 MIO WIG WIS
	50 55 60	
7.0		
70	too gae god gtd ato gtg ogo ago dog coa ggo ggo ba	c ttg ccc ggg 240
	Ser Asp Ala Val Ile Val Arg Ser Pro Pro Gly Gly Hi	s Leu Pro Gly
75	65 70 75	80
, 5		
	ccg cag cag caa gtg ccg ccg tcc cgc aac ggc tgt tc	c acc ctg ttt 288

- 23 -

	Pro	Gln	Gln	Gln	Val	Pro	Pro	Ser	Arg	Asn	Gly	Cys	Ser	Thr	Leu	Phe	
					85					90					95		
5																	
	agc	gac	atc	gct	ggc	gtc	aag	cđa	ctc	೩ಆಡ	GGG	gac	gat	tgg	ttg	gcc	336
10	Ser	Asp	Ile	Ala	cly	Val	Lys	Arg	Leu	Arģ	Pro	Авр	Asp	\mathtt{Trp}	Leu	Ala	
, Q				100					105					110			
15	gtc	aac	tcg	ccg	ccc	gcc	tct	tog	ccc	ತಿತಿದ	acg	tog	¢ac	ata	tcc	tac	384
	val	Asn	Ser	Pro	Pro	Ala	Ser	Ser	Pro	Gly	Thr	Ser	His	Ile	Ser	Tyr	
20			115					120					125				
20																	
	aca	gtc	ata	tcg	aac	99 ¢	99¢	ggc	ggt	9 90	ggc	ggt	ggc	33c	ggt	ggt	432
25	Thr	Val	Ile	Ser	Asn	Gly	Gly	Gly	Gly	Gly	Gly	Gly	Gly	Gly	Gly	Gly	
		130					135					140					
30																	
50	tac	aac	acg	tct	cca	atg	Ecg	acc	aac	agc	tac	gac	ccg	tac	agt	ccg	480
	Tyr	Asn	Thr	Ser	Pro	Met	Ser	Thr	Asn	Ser	Tyr	Asp	Pro	Tyr	Ser	Pro	
35	145					150					155					160	
40	atg	agt	gga	aaa	atc	gtc	aaa	gaa	gag	ttg	tct	ccg	cca	aac	agc	ctg	528
40	Met	Ser	Gly	Lys	Ile	Val	Lys	Glu	Glu	Leu	Ser	Pro	Pro	Asn	Ser	Leu	
					165					170					175		
45																	
	teg	gga	gtc	agc	agc	cat	tog	gat	333	ttg	aag	aag	aag	aaa	ctc	aac	576
50	Ser	Gly	Val	Ser	Ser	His	Ser	Asp	Gly	Ļeu	Lys	Lys	Lys	Lys	Leu	Asn	
30				180					185					190			
55	cac	ತರಡ	cc¢	tcg	acc	ggt	gtc	gtc	aac	acc	tcg	gca	tog	ggc	CCC	999	624
	His	Thr	Pro	Ser	Thr	GLy	Val	٧al	Asn	Thr	Ser	Ala	Ser	Gly	Pro	Gly	
60			195					200					205				
00																	
	ggt	33 c	gtt	ggt	ggc	aat	gtg	ctg	aac	aac	cga	cct	ccc	gaa	gag	ctg	672
65	Gly	Gly	Val	Gly	Gly	Asn	Val	Leu	Asn	Asn	Arg	Pro	Pro	Glu	Glu	Leu	
		210					215		•			220					
70																	
70	tgc	ctg	gtg	tgt	ggc	gac	cgg	tcg	tcc	₫g¢	tac	cat	tac	aac	gct	ctc	720
	Cys	Leu	Val	Сув	Gly	Asp	Arg	Ser	Ser	Gly	Тух	His	Tyr	Asn	Ala	Leu	
75	225					230					235					240	
	aca	tga	gaa	gga	tgc	aag	999	ttc	ttc	cgg	agg	agc	atc	acc	aag	aac	768
80	Thr	Cys	Glu	Gly	Cys	Lys	Gly	Phe	Phe	Arg	Arg	Ser	Ile	Thr	Lys	Asn	
		-		-	245	-	-			250	_				255		
85																	
	gcc	ਕੁਵਕ	tac	çag	tgc	aaq	tac	ggc	aac	aat	tge	gaa	ato	gac	atq	tac	816
	-			_	_	_					-	-		_	•	Tyr	
90			_	260	-	•	•	4	265					270			
				_													

- 24 -

5	_		cgg Arg	_	-	_		Cys					Cys				8 64
			275					280					285				
10	ggc	atg	agg	cct	gaa	tgt	gtt	gta	cct	gaa	gtt	çaa	tgc	gca	gta	aaa	912
	Gly	Met	Arg	Pro	Glu	Сув	Val	Val	Pro	Glu	Val	Gln	Суѕ	Ala	۷al	Lys	
15		290					295					300					
	292	aag	gag	aaa	aaa	gct	caa	cga	gaa	aaa	gat	aaa	cca	aat	tct	act	960
20	_	_	Glu													_	
	305					310					315					320	
25																	
		_	att			-					-				_	_	1008
30	THE	Asp	Ile	ser	325	GLU	TTE	T76	шyз	330	GIU	FLU	+ ****	G1.0	335	пÅъ	
										- "							
35	att	gaa	tgt	39¢	gaa	cca	atg	ata	atg	990	aca	cct	atg	cca	act	gta	1056
	Ile	Glu	сЛа	Gly	Glu	Pro	Met	Ίle	Met	Gly	Thr	Pro	Mct	Pro	Thr	Val	
40				340					345					350			
	aat	tar	gtq	222	aat	t.ta	agt	tet	graa	caa	ลลล	o a a	cto	atc	cac	coa	1104
45			Val			_	_		_			-	_			_	
		_	355	7				360			_		365				
50																	
		_	tat			_			_	- .		_	_			_	1152
55	Leu	Val 370	Tyr	Phe	Gln	Asp	Gln 375	Tyr	Glu	Ala	Pro	Ser 380	Glu	Lys	Asp	Met	
00		370					313					200					
60	aaa	cgt	tta	aca	ata	aat	aat	caa	aac	atg	gat	gaa	tat	gat	gaa	gaa	1200
00	Lys	Arg	Leu	Thr	Ile	Asn	Asn	Gln	Asn	Met	Asp	Glu	Tyr	Asp	Glu	Glu	
	385					390					395					400	
65	222		200	~~ ~	200	252	E		250	250		~~~	= + ~	200	222	ctc	1248
			Ser	_				_	_				_		_		1240
70	-1-				405		-1-	5		410				,	415		
75	aca	gtt	caa	ctg	att	gtt	gag	ttt	gcc	aaa	cga	tta	cca	ggt	tta	gat	1296
	Thr	Va <u>l</u>	Gln		11e	Val	Glu	Phe		•	Arg	Leu	Pro	_	Phe	Asp	
80				420					425					430			
	aaa	ctt	ata	aga	gaa	gat	caa	atc	act	tta	ctc	aaq	act	tac	tca	agt	1344
85				_	~	_						_	_	_		Ser	
			435	_				440					445	-			
90																	
- -	ģáa	gct	atg	atg	tto	agg	gta	gca	agg	aag	tat	gac	ato	acc	act	gac	1392

- 25 -

5	Glu	Ala 450	Met	Met	Phe	Arg	Val 455	Ala	Arg	Lys	Тут	Авр 460	Ile	Thr	Thr	Asp	
			gtg														1440
10	Ser 465	Ile	Val	Phe	Ala	Asn 470	Asn	Gln	Pro	Phe	Ser 475	Ala	Asp	ser	lyr	ASD 480	
	705																
15		_	gga	-													1488
	Lys	Ala	GJÀ	Leu	Gly 485	Asp	Ala	Ile	GIu	Asn 490	Gin	Leu	ser	Pne	ser 495	Arg	
20					103												
25		_	cac		_	_											1536
25	Phe	Met	Tyr	Asn 500	Met	Lys	Val	Asp	505	ATS	GIU	TYY	Ala	51 0	ren	Thr	
30																	
30	•		gtc			_	_							-			1584
35	Ala	Ile	Val 515	ITE	Pne	Ser	Ser	Arg 520	Pro	А\$П	ren	Leu	ASP 525	GIA	1.1.b	LYS	
40			aaa			-										_	1632
	vaı	530	Lys	11 6	GIN	GIU	535	Tyr	ren	GIU	ser	54 D	БУБ	WIG	TÄT	AGT	
45																	
	-		cga	_	_	_		_		_	_			_			1680
50	545	ASII	Arg	waħ	wrā	750	1115	Ala	THE	447	555	ıyı	VIC	~~3	<u> L</u> Gu	560	
55																cta Leu	1728
60	SET	AGT	nea	1111	565	nea	mag	TILL	Tea	570	ASII	914	. Adii		575	ыса	
60																	
65	-	_		_		_			_	_	_				-	Ala	1776
	СуБ	Med	1112	580	шyБ	NC C	шуа	, repii	585					590		113,0	
70																	
	-		rgg cgg	_	_	_											1797
<i>7</i> 5	014	110	595	_	744												
80	<21	0> 1	٥														
		1> 5															
85		.2> E															
	<21	.3 > N	lyzus	per	siça	tė											
90		_	_														
	<40) (>]	.0														

- 26 -

_	Met 1	Met	Asp	Gln	Lys 5	Cys	Asp	Val	Gly	Gly 10	Gly	Gly	Val	Ala	Ala 15	Ala
5	Ala	Ala	Gly	Ile 20	Gly	Gly	Gly	Gly	Val 25	Gly	Ģlу	Leu	Met	Ser 30	Tyr	Asn
10	Arg	Gly	Arg	Gly	Gly	Thr	Glu	Val	Ile	Ile	Lys	Pro	Arg	Ser	Pro	Ala
15			35					40					45			
20	Val	Val 50	Gln	Val	Ala	Thr	Gly 55	Gly	Ser	Tyr	His	60 Gly	Leu	Pro	Ala	Ala
25	Ser 65	Asp	Ala	Val	Ile	Val 70	Arg	Ser	Pro	Pro	Gly 75	Gly	His	Leu	Pro	Gly 80
30	Pro	Gln	Gln	Gln	Val 85	Pro	Pro	Şer	Arg	Asn 90	Gly	Сув	Ser	Thr	Leu 95	Phe
35	Ser	Asp	Ile		Gly	Val	Lys	Arg		Arģ	Pro	Asp	Asp	_	Leu	Ala
40				100	_			_	105				,	110		_
45	Val	Asn	Ser	Pro	Pro	Ala	Ser	Ser 120	Pro	Gly	Thr	Ser	125	Ile	ser	Tyr
50	Thr	Val 130	Ile	Ser	Asn	Gly	Gly 135	Gly	Gly	Gly	Gly	Gly 140	Gly	Gly	Gly	Gly
55	Tyr 145	Asn	The	Ser	Pro	Met 150	Ser	Thr	Asn	Ser	ïyr 155	Asp	Pro	Tyr	Ser	Pro 160
60	Met	Ser	Gly	Lys	Ile 165	V al	Lys	Glu	Glu	Leu 170		Pro	Pro	Asn	Ser 175	
65	Ser	Gly	Val	Ser	Ser	His	Ser	_Asp	Gly	Leu	Lys	Lys	Lys	Lys	Leu	Asn
70				180					185					190		
75	His	Thr	Pro 195		Thr	Gly	Val	. Val 200		Thr	Ser	Ala	Ser 205		Pro	Gly
80	Gly	Gly	v Val	Gly	Gly	Asn	Val 215		. Asn	Asn	Arg	220		Glu	. Glu	. Leu
85	Суя 225		. Val	. Cys	Gly	. Asp 230	_	, Ser	: Ser	Gly	7 Tyr 235		Tyr	Ası	a Ala	Leu 240
90	The	- ~u-	. Gl.	. Glv	r Cve	T.1/e	: C):	s Dhe	. Dhe	. 23 ≠c	, A-c	, Sei	- Tla	. ጥሎ ፣	- Taga	2 2 GT

- 27 -

					245					250					255	
5	Ala	Val	Tyr	Gln 260	Cys	Lys	Tyr	Gly	Asn 265	Asn	Cys	Glu	Ile	Asp 270	Met	Tyr
10	Met	Arg	Arg 275	Ľ у з	Cys	Gln	Glu	Cys 280	Arg	Leu	Lys	Lys	Cys 285	Leu	Thr	Val
15 20	Gly	Met 290	Arg	Pro	Glu	Cys	Val 295	Val	Pro	Glu	Val	Gln 300	Сув	Ala	Val	Lys
25	Arg	Lys	Gl u	Lys	Lys	Ala 310	Gln	Arg	Glu	L y ≉	Авр 315	Lys	Pro	Asn	Ser	Thr 320
30	Thr	Asp	Ile	Ser	Pro 325	Glu	Ile	Ile	Lys	11e	Glu	Pro	Thr	Glų	Met 335	Lys
35	Ile	G lu	Cys	Gly 340	Glu	Pro	Met	Ile	Met 345	Gly	Thr	Pro	Met	Pro 350	Thr	Val
40	Pro	Tyr	Val 355	Lys	Pro	Leu	Ser	Ser 360	Glu	Gln	Lys	Glu	Leu 365	Ile	His	Arg
45 50	Leu	Val	Tyr	Phe	Gln	Asp	Gln 375	Tyr	Glu	Ala	Pro	Ser 380	Glu	Lys	Asp	Met
55	L ys 385	Arg	Leu	Thr	Ile	Asn 390	Asn	Gln	Asn	Met	Asp 395	Glu	тут	Asp	Glu	Glu 400
60	Lys	Gln	Ser	Asp	Thr 405	Thr	Tyr	Arg	Ile	Ile	Thr	Glu	Met	The	Ile 415	Γ¢π
65	Thr	Val	Gln	Leu 420	Ile	Val	Glu	Phe	Ala 425	Lys	Arg	Leu	Pro	Gly 430	Phe	Asp
70	Lys	Leu	Val 435	Arg	Glu	Asp	Gln	Ile 440		Leu	Leu	Lys	Ala		Ser	Ser
75 80	Glu	Ala 450		Met	Phe	Arg	Val 455		Arg	Lys	Tyr	Asp	Ile	Thr	Thr	Авр
85	Ser 465		val	Phe	Ala	Asn 470		. Gln	Pro	Phe	Ser 475		Asp	Sez	Tyz	Asn 480
90	L y s	Ala	. Gly	- Leu	1 Gly 485		Ala	ı Ile	e Glu	490		ı Leu	. Sei	r Phe	Ser 495	Arg

- 28 -

5	Phe I	Met	Tyr	Asn 500	Met	Lys	Val	Asp	Aşn 505	Ala	Glu	Tyr	Ala	Leu 510	Leu	Thr	
10	Ala :	Ile	val 5 1 5	Ile	Phe	Ser	Ser	Arg 520	Pro	Asn	Leu	Leu	Asp 525	Gly	Trp	Lys	
15	Val (Glu 530	Lys	Ile	Gln	Glu	Ile 535	Тут	Leu	Glu	Ser	Leu 5 40	Lys	Ala	Tyr	Val	
20	Asp 5	Asn	Arg	qeA	Arg	Asp 550	Thr	Ala	Thr	Val	A rg 555	Tyr	Ala	Arg	Leu	Leu 560	
25 30	Ser '	Val	Leu	Thr	Glu 565	Leu	Arg	Thr	Leu	Gly 570	Asn	Glu	Asn	Ser	Glu 575	Leu	
35	Cys)	Met	Thr	Leu 580	Lys	Leu	Lys	Asn	Arg 585	Val	Val	Pro	Pro	Phe 590	Leu	Ala	
40	Glu	Ile	Trp 595	Азр	Val	Met	Pro										
45	<210	> 13	Ĺ														
50	<211 <212																
<i>5</i> 5	<213 <220 <221	>		per	sica	Ė											
60	<222	> (]	L)	(113	L)												
65	<400 atg Met	tat	tcc				_								tac Tyr		4 B
70	1				5					10					15		
75	_	-				_	_				_		-		gac Asp		96
80	_			-						-					ctg	_	144
85	PIO	asn	35		PTO	ASN	nis	40	ьeu	zer	чТĀ	ser	45		Геп	сув	
90			_		_	_	_	-						_		agc Ser	192

- 29 -

tge gag ggg tge aaa ggg tte tte aaa cgc aca gtg agg aaa aat ttg Cys Glu Gly Cys Lys Gly Phe Phe Lys Arg Thr Val Arg Lys Asn Leu tea tad gog tgt ege gaa gaa aac aaa tgc atc atc gac aag ege caa Ser Tyr Ala Cys Arg Glu Glu Asn Lys Cys Ile Ile Asp Lys Arg Gln oga aat ogg tge caa tac tge agg tat caa aaa tgt ttg acc atg gge Arg Asn Arg Cys Gln Tyr Cys Arg Tyr Gln Lys Cys Leu Thr Met Gly atg ann aga gaa gct gtg cag gan gan aga can cgt aca naa gan cga Met Lys Arg Glu Ala Val Gln Glu Glu Arg Gln Arg Thr Lys Glu Arg gat cat aat aac atc gaa gtt gaa coc acg agc agt tot aat act gat Asp His Asn Asn Ile Glu Val Glu Pro Thr Ser Ser Asn Thr Asp atg cca gtg gaa ctc ata tta agg gct gag aat aaa gct gat gct ata Met Pro Val Glu Leu Ile Leu Arg Ala Glu Asn Lys Ala Asp Ala Ile aag act gaa caa cag tat ata gag caa cga cat cct caa cat act gtt Lys Thr Glu Gln Gln Tyr Ile Glu Gln Arg His Pro Gln His Thr Val ggt gct att tgt caa gca act gac aag cag tta ata caa ctt gtt gaa Gly Ala Ile Cys Gln Ala Thr Asp Lys Gln Leu Ile Gln Leu Val Glu 1.85 tgg gcc aag cat ata ccg cat ttt aaa aat tta cct cta ggc gat caa Trp Ala Lys His Ile Pro His Phe Lys Asn Leu Pro Leu Gly Asp Gln gtt tta tta ttg aga gct ggt tgg aat gag ttg atg att gca gca ttt Val Leu Leu Arg Ala Gly Trp Asn Glu Leu Met Ile Ala Ala Phe too cat aga toa atc agt gta aaa gat ggt ata gto tta gct act gga Ser His Arg Ser Ile Ser Val Lys Asp Gly Ile Val Leu Ala Thr Gly

- 30 -

	ctt	act	gtt	gac	aga	gat	tca	gct	cac	caa	ācŗ	ggt	gţţ	gaa	gct	ata	768
	Leu	Thr	Val	Asp	Arg	Asp	Ser	Ala	His	Gln	Ala	Gly	Val	Glu	Ala	Ile	
5					245					250					255		
	+++	cat	rat	ota	ctc	act	gaa	ctc	att	art	222	ata	aga	gat	atg	aat	816
10		_	_	_			_		•	_		_	_	-	Met		
		1.175PA	74.9		110.0	1111	GIG	пец		ALG	шу.э	MCC	A	-	1.14	C.L.y	
15				260					265					270			
15																	
	_	_	_					_	_	_					CEE		864
20	Met	Asp	Arg	Thr	Glu	Leu	Gly	Cys	Leu	Arg	The	Ile	Ile	Leu	Phe	Asn	
			275					280					285				
25	ÇÇZ	ggt	tca	aaa	ggt	ttg	cag	tct	gtg	aat	gaa	gtg	caa	gta	ctg	cgt	912
	Pro	Gly	Ser	Lys	Gly	Leu	Gln	Ser	Val	Asn	Glu	Val	Gln	Val	Leu	Arg	
20		290					295					300					
30																	
	gat	aag	gtt	tat	gtt	geg	tta	qaa	qaa	tat	tat	çgt	aca	aca	cat	CCA	960
35											-				His		
	305			- 3		310				-1-	315	د				320	
																220	
40		~~ n															7.000
					_										tca		1008
45	GIU	GIU	PEO	GIA	_	Fre	ATa	гуя	Leu		Leu	Arg	Leu	PIO	Ser	Leu	
45					325					330					335		
50	cgt	tca	act	ààs	tta	aaa	tgt	ctg	gaa	çat	tta	ttc	ttt	tat	aaa	ctt	1056
	Arg	Ser	Ile	Ģlγ	Leu	Lys	Cys	Leu	Glu	His	Leu	Phe	Phe	Tyz	Lys	Leu	
				340					345					350			
<i>5</i> 5																	
	att	ggc	gat	tcc	cca	att	gat	aca	ttt	t ta	atg	gaa	gtt	ctc	gaa	tca	1104
	_														Glu		
60		_	355				•	360					365				
65	r at	tas	oa F	gac	~++	~39	7 = 0	~~+	107								1131
-				_	_		_	_									1121
	Ser		TTE	Asp	Val	GIII		wid	Int								
70		370					375										
75	<21	0> 1	2														
	<21	1 > 3	77														
80	<21	2> P	RT														
UU	<21	3 > M	yzus	per	sica	e											
85																	
		0> 1															
	Met	Tyr	Ser	Asn	Ser	Tyr	Thr	Met	Tyr	Ser	Ser	Asp	Arg	Leu	Tyr	Ser	
90	1				5					10					15		

- 31 -

5	Val	Asp	Arg	Asn 20	Ser	Met	Met	Asn	Asn 25	Ser	Cys	Asn	Val	Gln 30	Авр	Ser
	Pro	Asn	Tyr 35	Pro	Pro	Asn	His	Pro 40	Leu	Ser	Gly	Ser	Lys 45	His	Leu	Сув
0	Ser	αľΤ		GIV	Δ«n	Ara	ala		G) v	Tars	Hi a	TV ァ	Gly	Val	Tvr	Ser
5		50	-7.5	,		3	55		 1	-,-		60			-1-	
20	Çy≉ 65	Glu	Gly	Cys	Lys	Gly 70	Phe	Phe	ŗÀa	Arg	Thr 75	Val	Arg	Lys	Asn	Leu so
25	Ser	Tyr	Ala	Сув	Arg 85	Glu	Glu	Asn	Lys	Cys 90	Ile	Ile	Asp	Lys	Arg 95	Gln
30 35	Arg	Asn	Arg	Cys 100	Gln	Tyr	Cys	Arg	Tyr 105	Gln	Lys	Сув	Leu	Thr 110	Met	Gly
10	Met	Lys	Arg 115	Glu	Ala	Val	Gln	Glu 120	Glu	Arg	Gln	Arg	Thr 125	Lys	Glu	Arg
\$5	Asp	His 130	Asn	Aŝn	Ile	Glu	Val 135	Glu	Pro	Thr	Ser	Ser 140	Ser	Asn	Thr	Asp
50	Met 145	Pro	Val	Glu	Leu	Ile 150	Leu	Arg	Ala	Glu	Asn 155	Lys	Ala	Asp	Ala	Ile 160
55	Lys	Thr	Glu	Gln	Gln 165	Tyr	Ile	Glu	Gln	Arg 170	His	Pro	Gln	His	Th r 175	Val
60 	Gly	Ala	Ile	Cys	Gln	Ala	Thr	Asp	Lys 185	Gln	Leu	Ile	Gln	Leu 190	Val	Glu
55 70	Trp	Ala	L уs 195	His	ĭle	Pro	His	Phe 200	īys	Asn	Leu	Pro	Leu 205	Gly	Asp	Gln
75	Val	Leu 210	Leu	Leu	Arg	Ala	Gly 215	Trp	Asn	Glu	Leu	Met 220	Ile	Ala	Ala	Phe
80	Ser 225	His	Arg	Ser	Ile	Ser 230	Val	Lys	Asp	Gly	Ile 235	Val	Leu	Ala	Thr	Gly 240
85	Lėų	Thr	Val	Asp	Arg 245	Asp	Ser	Ala	His	Gln 250		Gly	Val	Glu	Ala 255	Ile
9 0	Phe	Asn	Arc	Val	7. 0 11	Th-	יי לם	Len	บรไ	בומ	Larg	Mor	Ara	ገ ረም	Met	c1.

- 32 -

260 265 270 Met Asp Arg Thr Glu Leu Gly Cys Leu Arg Thr Ile Ile Leu Phe Asn 280 10 Pro Gly Ser Lys Gly Leu Gln Ser Val Asn Glu Val Gln Val Leu Arg 295 15 Asp Lys Val Tyr Val Ala Leu Glu Glu Tyr Cys Arg Thr Thr His Pro 305 310 315 20 Glu Glu Pro Gly Arg Phe Ala Lys Leu Leu Arg Leu Pro Ser Leu 25 325 Arg Ser Ile Gly Leu Lys Cys Leu Glu His Leu Phe Phe Tyr Lys Leu 30 340 345 35 Ile Gly Asp Ser Pro Ile Asp Thr Phe Leu Met Glu Val Leu Glu Ser 360 40 Ser Ser His Asp Val Gln Val Ala Thr 370 375 45 <210> 13 <211> 150 <212> DNA 50 <213> Lucilia cuprina <220> <221> CDS <222> (9)..(134) 55 <400> 13 aattetge gaa gga tge aag gga tte tte aaa egt ace gta ege aag gae 50 Glu Gly Cys Lys Gly Phe Phe Lys Arg Thr Val Arg Lys Asp

ttg aca tat gct tgt cgt gag gac aga aat tgc att ata gat aaa cga 98

10

5

1

60

- 33 -

Leu Thr Tyr Ala Cys Arg Glu Asp Arg Asn Cys Ile Ile Asp Lys Arg

15

20

25

30

caa aga aat ogt tgo cag tat tgt ogo tao caa aag tgatogatac ogtoga 150

5 Gln Arg Asn Arg Cys Gln Tyr Cys Arg Tyr Gln Lys

35

40

<210> 14

10 <211> 42

<212> PRT

<213> Lucilia cuprina

<400> 14

Glu Gly Cys Lys Gly Phe Phe Lys Arg Thr Val Arg Lys Asp Leu Thr

15

5

10

15

Tyr Ala Cys Arg Glu Asp Arg Asn Cys Ile Ile Asp Lys Arg Gln Arg

20

25

30

20 Asn Arg Cys Gln Tyr Cys Arg Tyr Gln Lys

35

40

<210> 15

25 <211> 32

<212> DNA

<213> Artificial Sequence

<220>

30 <223> Description of Artificial Sequence:primer

5

- 34 -

<400> 15			
cggaattccg	cctcnggnta	ycaytayaay	gc

32

```
<211> 32
<212> DNA
<213> Artificial Sequence
```

<210> 16

<210> 17

32

15

20

<211> 23
<212> DNA
<213> Artificial Sequence
<220>

<223> Description of Artificial Sequence:primer

25 <400> 17
gcctcggggt atcactataa cgc

23

<210> 18

30 <211> 23

<212> DNA

10

<210> 19

- 35 -

	<213> Artificial Sequence
	<220>
	<223> Description of Artificial Sequence:primer
5	
	<400> 18
	gcacteetga cactttegte tea

23

```
<211> 23
     <212> DNA
     <213> Artificial Sequence
15
     <220>
     <223> Description of Artificial Sequence:primer
     <400> 19
20
    tegteeggtt accattacaa ege
```

<210> 20 <211> 24 25 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:primer 30 <400> 20

tagacetttg geraaytena caat

23

24

JOINT INVENTORS' DECLARATION FOR PATENT APPLICATION AND POWER OF ATTORNEY

As the below named inventors, we hereby declare that:

Our residences, post office addresses and citizenship are as stated below our names.

We believe that we are the original, first and joint inventors of the subject matter which is claimed and for which a patent is sought on the invention entitled: "Genetic Sequences Encoding Steroid and Juvenile Hormone Receptor Polypeptides and Insecticidal Modalities Therefor," the specification of which was filed on 1 July 1999, as Application Serial No.

We hereby authorize our legal representative to add reference to the Serial No. and/or filing date of the above-referenced application to this declaration.

We hereby state that we have reviewed and understand the contents of the aboveidentified specification, including the claims, as amended by any amendment referred to above.

We acknowledge the duty to disclose information which is material to the patentability of this application in accordance with Title 37, Code of Federal Regulations, §1.56.

Prior Foreign Application(s)

We hereby claim foreign priority benefits under Title 35, United States Code, §119 of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application(s) for patent or inventor's certificate having a filing date before that of the application to which priority is claimed:

Country	Application No.	Date of Filing (day,month,year)	Date of Issue (day,month,year)	Priority Claimed 35 U.S.C.119	
AU	PP1356/98	15 January 1999		Yes <u>X</u> No	

Prior Provisional Application(s)

We hereby claim the benefit under Title 35, United States Code, §119(e) of any United States provisional application(s) listed below:

Application Serial Number Date of Filing (day,month,year)

Prior U.S. Application(s) and PCT International Application(s) Designating the United States

We hereby claim the benefit under Title 35, United States Code, §120 of any United States application(s), or § 365(c) of any PCT International application(s) designating the United States listed below:

Application Serial

Date of Filing

Status(Patented, Pending, Abandoned)

Number

(day,month,year)

PCT/AU99/00033

15 January 1999

Pending

Insofar as the subject matter of each of the claims in this application is not disclosed in the prior United States, foreign or PCT International application(s) to which priority has been claimed above in the manner provided by the first paragraph of Title 35, United States Code, §112, we acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, §1.56 which occurred between the filing date of the prior application(s) and the national or PCT international filing date of this application.

We hereby appoint, both jointly and severally, as our attorneys and agents with full power of substitution and revocation, to prosecute this application and any corresponding application filed in the Patent Cooperation Treaty Receiving Office, and to transact all business in the Patent and Trademark Office connected herewith the following attorneys and agents, their registration numbers being listed after their names:

Lorance L. Greenlee, Reg. No. 27,894; Ellen P. Winner, Reg. No. 28,547; Sally A. Sullivan, Reg. No. 32,064; Donna M. Ferber, Reg. No. 33,878; No. 37,374; G. William VanCleave, Reg. No. 40,213; and Susan K. Doughty, Reg. No. 43,595, all of Greenlee,

(1) Full Name of

Winner and Sullivan, P.C., 5370 Manhattan Circle, Suite 201, Boulder, CO 80303.

We hereby declare that all statements made herein of our own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

First Inventor: Residence:	HILL, Ronald Johnston Forestville, New South Wales, 2087, Australia
Citizenship:	Australia
Post Office Address:	Altona Avenue, Forestville, New South Wales, 2087, Australia
(1) Signature	Date
(2) Full Name of Second Inventor: Residence: Citizenship: Post Office Address:	HANNAN, Garry Noel Hunters Hill, New South Wales, 2110, Australia Australia 32 Earl Street, Hunters Hill, New South Wales, 2110, Australia
(2) Signature	Date